EL 617 042 690 US Express Mail Label No. August 24, 2001

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APPLICATION

for

UNITED STATES LETTERS PATENT

on

STRESS-REGULATED GENES OF PLANTS, TRANSGENIC PLANTS CONTAINING SAME, AND METHODS OF USE

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Sheets of Drawings: None Docket No.: SCRIP1300-3

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STRESS-REGULATED GENES OF PLANTS, TRANSGENIC PLANTS CONTAINING SAME, AND METHODS OF USE

[0001] This application claims the benefit under 35 U.S.C. 119(e) of U.S. Serial No. 60/227,866, filed August 24, 2000; U.S. Serial No. 60/264,647, filed January 26, 2001; and U.S. Serial No. 60/300,111, filed June 22, 2001, each of which is incorporated herein by reference in its entirety.

[0002] Three CD-R compact discs, labeled "Copy 1", "Copy 2", and "CRF" and having the files listed below, are submitted herewith and are incorporated herein by reference. Copy 1 and Copy 2 each contain two text documents: 1) a file named SCRIP1300-3_SEQUENCE_LISTING, which contains the Sequence Listing, was created on August 20, 2001 (and recorded on the CD-R on August 21, 2001), and is 9,972 KB in size; and 2) a file named SCRIP1300-3_Table_32, which contains Table 32, was created on August 20, 2001 (and recorded on the CD-R on August 21, 2001), and is 1,251 KB in size. The CRF contains a single file named SCRIP1300-3_SEQUENCE_LISTING, which contains the Sequence Listing, was created on August 20, 2001 (and recorded on the CD-R on August 21, 2001), is 9,972 KB in size, and is identical to the files having the same name on Copy 1 and Copy 2.

BACKGROUND OF THE INVENTION FIELD OF THE INVENTION

[0003] The present invention relates generally to plant genes, the expression of which are regulated in response to stress, and more specifically to the gene regulatory elements involved in a stress-induced response in plants, to uses of the coding sequences and regulatory elements of such plant stress-regulated genes, and to transgenic plants genetically modified to express such a coding sequence or to express a heterologous polynucleotide from such a regulatory element.

BACKGROUND INFORMATION

[0004] Microarray technology is a powerful tool that can be used to identify the presence and level of expression of a large number of polynucleotides in a single assay. A microarray is formed by linking a large number of discrete polynucleotide sequences, for example, a population of polynucleotides representative of a genome of an organism, to a solid support such as a microchip, glass slide, or the like, in a defined pattern. By contacting the microarray with a nucleic acid sample obtained from a cell of interest, and detecting those polynucleotides expressed in the cell can hybridize specifically to complementary sequences on the chip, the pattern formed by the hybridizing polynucleotides allows the identification of clusters of genes that are expressed in the cell. Furthermore, where each polynucleotide linked to the solid support is known, the identity of the hybridizing sequences from the nucleic acid sample can be identified.

[0005] A strength of microarray technology is that it allows the identification of differential gene expression simply by comparing patterns of hybridization. For example, by comparing the hybridization pattern of nucleic acid molecules obtained from cells of an individual suffering from a disease with the nucleic acids obtained from the corresponding cells of a healthy individual, genes that are differentially expressed can be identified. The identification of such differentially expressed genes provides a means to identify new genes, and can provide insight as to the etiology of a disease.

[0006] Microarray technology has been widely used to identify patterns of gene expression associated with particular stages of development or of disease conditions in animal model systems, and is being applied to the identification of specific patterns of gene expression in humans. The recent availability of information for the genomes of plants provides a means to adapt microarray technology to the study of plant gene expression.

[0007] Plants and plant products provide the primary sustenance, either directly or indirectly, for all animal life, including humans. For the majority of the world's human population and for many animals, plants and plant products provide the sole source of nutrition. As the world population increases, the best hope to prevent widespread famine is to increase the quantity and improve the quality of food crops, and to make the crops available to the regions of the world most in need of food.

[0008] Throughout history, a continual effort has been made to increase the yield and nutritious value of food crops. For centuries, plants having desirable characteristics such as greater resistance to drought conditions or increased size of fruit were crossbred and progeny plants exhibiting the desired characteristics were selected and used to produce seed or cuttings for propagation. Using such classical genetic methods, plants having, for example, greater disease resistance, increased yield, and better flavor have been obtained. The identification of plant genes involved in conferring a selective advantage on the plant to an environmental challenge would facilitate the generation and yield of plants, thereby increasing the available food supply to an increasing world population. The involvement of these genes in a single organism to responses to multiple stress conditions, however, remains unknown.

Thus, a need exists to identify plant genes and polynucleotides that are involved in modulating the response of a plant to changing environmental conditions. The present invention satisfies this need and provides additional advantages.

SUMMARY OF THE INVENTION

[0009] The present invention relates to clusters of genes that are regulated in response to a stress condition in plants. Such clusters include, for example, plant polynucleotides whose expression is altered in response to two or more different stress conditions; and plant polynucleotides the expression of which are altered in response to one stress condition, but not to others. The identification of such clusters, using microarray technology, has allowed the identification of plant stress-regulated genes in *Arabidopsis thaliana* (see Tables 1 and 2); and homologs and orthologs thereof in other plant species (see Table 32). Thus, the invention provides isolated

polynucleotide portions of *Arabidopsis* plant stress-regulated genes, and homologs and orthologs thereof; variants of such sequences, and polynucleotides encoding substantially similar plant stress-regulated polypeptides expressed therefrom. Such sequences include, for example, sequences encoding transcription factors; enzymes, including kinases; and structural proteins, including channel proteins (see Tables 29-31). Accordingly, the present invention also relates to an isolated polynucleotide comprising all or a portion of a plant stress-regulated gene, and to polynucleotide portions thereof, including a coding region (open reading frame), which encodes all or a portion of a stress-regulated polypeptide, for example, as set forth in SEQ ID NOS:1-2703; and a regulatory element involved in regulating the response of the plant to a stress condition such exposure to an abnormal level of salt, osmotic pressure, temperature or any combination thereof, for example, as set forth in SEO ID NOS:2704-5379.

[0010] The present invention also relates to a recombinant polynucleotide, which contains a nucleotide sequence of a plant stress-regulated gene or functional portion thereof operatively linked to a heterologous nucleotide sequence. In one embodiment, the recombinant polynucleotide comprises a plant stress-regulated gene regulatory element operatively linked to a heterologous nucleotide sequence, which is not regulated by the regulatory element in a naturally occurring plant. The heterologous nucleotide sequence, when expressed from the regulatory element, can confer a desirable phenotype to a plant cell containing the recombinant polynucleotide. In another embodiment, the recombinant polynucleotide comprises a coding region, or portion thereof, of a plant stress-regulated gene operatively linked to a heterologous promoter. The heterologous promoter provides a means to express an encoded stress-regulated polypeptide constitutively, or in a tissue-specific or phase-specific manner.

[0011] Accordingly, in one aspect, the present invention provides an isolated polynucleotide comprising a nucleotide sequence of a plant gene that hybridizes under stringent conditions, preferably high stringency conditions, to any one of SEQ ID NOS:1-5379 (see Tables 1 and 2), including to a coding region (SEQ ID

NOS:1-2703) or a regulatory region, which can alter transcription of an operatively linked nucleic acid sequence in response to an abiotic stress (SEQ ID NOS:2704-5379; see Table 2), or to a complement thereof. Additional aspects provide sequences that hybridize under stringent conditions, preferably high stringency conditions, to the complements of SEQ ID NO 1-1261 (cold responsive genes; Tables 3-6), SEQ ID NOS:2227-2427 (saline responsive genes; Tables 7-10), SEQ ID NOS:2428-2585 (osmotic responsive genes; Tables 11-14), SEQ ID NOS:1699-1969 (cold and osmotic responsive genes; Tables 15-17), SEQ ID NOS:1970-2226 (cold and saline responsive genes; Tables 18-20), SEQ ID NOS:2586-2703 (osmotic and saline responsive genes; Tables 21-23), and SEQ ID NOS:1262-1698(cold, osmotic and saline responsive genes; Tables 24-26), and which can comprise regulatory regions that can alter transcription in response to cold stress, osmotic stress, saline stress, or combinations thereof (SEQ ID NOS:2704-5379; see Table 2). Also provided are nucleotide sequences complementary thereto, and expression cassettes, plants and seeds comprising any of the above isolated sequences.

[0012] In another aspect, the present invention provides an isolated polynucleotide comprising a plant nucleotide sequence that hybridizes under stringent conditions, preferably high stringency conditions, to the complement of any one of SEQ ID NOS:1-2703 (Table 1), including to a coding region thereof (SEQ ID NOS:2704-5379), wherein expression of said coding region is altered in response to an abiotic stress. Additional aspects provide sequences that hybridize under high stringency conditions to the complements of SEQ ID NO 1-1261 (cold responsive genes; Tables 3-6), SEQ ID NOS:2227-2427 (saline responsive genes; Tables 7-10), SEQ ID NOS:2428-2585 (osmotic responsive genes; Tables 11-14), SEQ ID NOS:1699-1969 (cold and osmotic responsive genes; Tables 15-17), SEQ ID NOS:1970-2226 (cold and saline responsive genes; Tables 18-20), SEQ ID NOS:1262-1698(cold, osmotic and saline responsive genes; Tables 21-23), and SEQ ID NOS:1262-1698(cold, osmotic and saline responsive genes; Tables 24-26), and which can comprise a coding region whose transcription is altered in response to cold stress, osmotic stress, saline stress, or a combination thereof. Also provided are nucleotide

sequences complementary thereto, and expression cassettes, plants and seeds comprising any of the above sequences.

[0013] The invention further relates to a method of producing a transgenic plant, which comprises at least one plant cell that exhibits altered responsiveness to a stress condition. In one embodiment, the method can be performed by introducing a polynucleotide portion of plant stress-regulated gene into a plant cell genome, whereby the polynucleotide portion of the plant stress-regulated gene modulates a response of the plant cell to a stress condition.

The polynucleotide portion of the plant stress-regulated gene can encode a stress-regulated polypeptide or functional peptide portion thereof (see SEO ID NOS:1-2703), wherein expression of the stress-regulated polypeptide or functional pentide portion thereof either increases the stress tolerance of the transgenic plant, or decreases the stress tolerance of the transgenic plant. The polynucleotide portion of the plant stress-regulated gene encoding the stress-regulated polypeptide or functional peptide portion thereof can be operatively linked to a heterologous promoter. The polynucleotide portion of the plant stress-regulated gene also can comprise a stressregulated gene regulatory element (see SEQ ID NOS:2704-5379). The stressregulated gene regulatory element can integrate into the plant cell genome in a sitespecific manner, whereupon it can be operatively linked to a heterologous nucleotide sequence, which can be expressed in response to a stress condition specific for the regulatory element; or can be a mutant regulatory element, which is not responsive to the stress condition, whereby upon integrating into the plant cell genome, the mutant regulatory element disrupts an endogenous stress-regulated regulatory element of a plant stress-regulated gene, thereby altering the responsiveness of the plant stressregulated gene to the stress condition.

[0015] In one aspect, the invention provides a method for producing a transgenic plant by introducing into at least one plant cell a recombinant nucleic acid construct comprising i) all or a portion of any one of SEO ID NOS:1-5379; ii) a polynucleotide

comprising a coding region that hybridizes under conditions of high stringency to all or a portion of the complement of any one of SEQ ID NOS:1-2703; iii) a polynucleotide comprising a sequence that alters transcription of an operatively linked coding region in response to abiotic stress, and that hybridizes under conditions of high stringency to the complement of any one of SEQ ID NOS:2704-5379; iv) a polynucleotide having at least 90% sequence identity with any one of SEQ ID NO:1-5379; v) a fragment of any one of the sequences of iv), wherein the fragment comprises a coding region; or vi) a fragment of any one of the sequences of iv), wherein the fragment comprises a nucleotide sequence that alters transcription of an operatively linked coding region in response to abiotic stress; and regenerating a plant from the at least one plant cell.

[0016] Another aspect provides a method for producing a transgenic plant comprising introducing into at least one plant cell a recombinant nucleic acid construct comprising i) any one of SEQ ID NOS:1-1261 or 2704-3955; ii) a polynucleotide comprising a coding region that hybridizes under conditions of high stringency to the complement of any one of SEQ ID NOS:1-1261; iii) a polynucleotide comprising a sequence that alters transcription of an operatively linked coding region in response to cold stress that hybridizes under conditions of high stringency to the complement of any one of SEQ ID NOS:2704-3955; iv) a polynucleotide that has at least 90% sequence identity with any one of SEQ ID NOS:1-1261 or 2704-3955; v) a fragment of any one of the sequences of iv), wherein the fragment comprises a coding region; or vi) a fragment of any one of the sequences of iv) wherein the fragment comprises a sequence or region that alters transcription of an operatively linked coding region in response to cold stress; and regenerating a plant from the at least one plant cell.

[0017] In another aspect, the invention provides a method for producing a transgenic plant by introducing into at least one plant cell a recombinant nucleic acid construct comprising i) any one of SEQ ID NOS:2428-2585 or 5108-5263; ii) a polynucleotide comprising a coding region that hybridizes under conditions of high

stringency to the complement of any one of SEQ ID NOS:2428-2585; iii) a polynucleotide comprising a sequence that alters transcription of an operatively linked coding region in response to osmotic stress that hybridizes under conditions of high stringency to the complement of any one of SEQ ID NOS:5108-5263; iv) a polynucleotide that has at least 90% sequence identity with any one of SEQ ID NOS:2428-2585 or 5108-5263; v) a fragment of any one of the sequences of iv), wherein the fragment comprises a coding region; or vi) a fragment of any one of the sequences of iv), wherein the fragment comprises a sequence or region that alters transcription of an operatively linked coding region in response to osmotic stress; and regenerating a plant from the at least one plant cell.

[0018] Still another aspect provides a method for producing a transgenic plant comprising introducing into at least one plant cell a recombinant nucleic acid construct comprising i) any one of SEQ ID NOS:2227-2427 or 4910-5107; ii) a polynucleotide comprising a coding region that hybridizes under conditions of high stringency to the complement of any one of SEQ ID NOS:2227-2427; iii) a polynucleotide comprising a sequence that alters transcription of an operatively linked coding region in response to saline stress that hybridizes under conditions of high stringency to the complement of any one of SEQ ID NOS:2227-2427; iv) a polynucleotide that has at least 90% sequence identity with any one of SEQ ID NOS:4910-5107; v) a fragment of any one of the sequences of iv) wherein the fragment comprises a coding region; or vi) a fragment of any one of the sequences of iv) wherein the fragment comprises a sequence or region that alters transcription of an operatively linked coding region in response to saline stress; and regenerating a plant from the at least one plant cell.

[0019] Yet another aspect provides a method for producing a transgenic plant comprising introducing into at least one plant cell a recombinant nucleic acid construct comprising i) any one of SEQ ID NOS:1699-1969 or 4389-4654; ii) a polynucleotide comprising a coding region that hybridizes under conditions of high stringency to the complement of any one of SEQ ID NOS:1699-1969; iii) a

polynucleotide comprising a sequence that alters transcription of an operatively linked coding region in response to a combination of cold and osmotic stress that hybridizes under conditions of high stringency to the complement of any one of SEQ ID NOS:4389-4654; iv) a polynucleotide that has at least 90% sequence identity with any one of SEQ ID NOS:1699-1969 or 4389-4654; v) a fragment of any one of the sequences of iv), wherein the fragment comprises a coding region; or vi) a fragment of any one of the sequences of iv), wherein the fragment comprises a sequence or region that alters transcription of an operatively linked coding region in response to a combination of cold and osmotic stress; and regenerating a plant from the at least one plant cell.

[0020] Yet another aspect provides a method for producing a transgenic plant comprising introducing into at least one plant cell a recombinant nucleic acid construct comprising i) any one of SEO ID NOS:1970-2226 or 4655-4909; ii) a polynucleotide comprising a coding region that hybridizes under conditions of high stringency to the complement of any one of SEO ID NOS:1970-2226; iii) a polynucleotide comprising a sequence that alters transcription of an operatively linked coding region in response to a combination of cold and saline stress that hybridizes under conditions of high stringency to the complement of any one of SEQ ID NOS:4655-4909; iv) a polynucleotide that has at least 90% sequence identity with any one of SEQ ID NOS:1970-2226 or 4655-4909; v) a fragment of any one of the sequences of iv), wherein the fragment comprises a coding region; or vi) a fragment of any one of the sequences of iv), wherein the fragment comprises a sequence or region that alters transcription of an operatively linked coding region in response to a combination of cold and saline stress; and regenerating a plant from the at least one plant cell.

[0021] A further aspect provides a method for producing a transgenic plant comprising introducing into at least one plant cell a recombinant nucleic acid construct comprising i) any one of SEQ ID NOS:2586-2703 or 5264-5379; ii) a polynucleotide comprising a coding region that hybridizes under conditions of high

stringency to the complement of any one of SEQ ID NOS:2586-2703; iii) a polynucleotide comprising a sequence that alters transcription of an operatively linked coding region in response to a combination of osmotic and saline stress that hybridizes under conditions of high stringency to the complement of any one of SEQ ID NOS: 5264-5379; iv) a polynucleotide that has at least 90% sequence identity with any one of SEQ ID NOS:2586-2703 or 5264-5379; v) a fragment of any one of the sequences of iv), wherein the fragment comprises a coding region; or vi) a fragment of any one of the sequences of iv), wherein the fragment comprises a sequence or region that alters transcription of an operatively linked coding region in response to a combination of osmotic and saline stress; and regenerating a plant from the at least one plant cell.

[0022] Another aspect provides a method for producing a transgenic plant comprising introducing into at least one plant cell a recombinant nucleic acid construct comprising i) any one of SEO ID NOS:1262-1698 or 3956-4388; ii) a polynucleotide comprising a coding region that hybridizes under conditions of high stringency to the complement of any one of SEQ ID NOS:1262-1698; iii) a polynucleotide comprising a sequence that alters transcription of an operatively linked coding region in response to a combination of cold, osmotic and saline stress that hybridizes under conditions of high stringency to the complement of any one of SEO ID NOS:3956-4388; iv) a polynucleotide that has at least 90% sequence identity with any one of SEO ID NOS:1262-1698 or 3956-4388; v) a fragment of any one of the sequences of iv), wherein the fragment comprises a coding region; or vi) a fragment of any one of the sequences of iv) wherein the fragment comprises a sequence or region that alters transcription of an operatively linked coding region in response to a combination of cold, osmotic and saline stress; and regenerating a plant from the at least one plant cell. Further aspects include plants and uniform populations of plants made by the above methods as well as seeds and progeny from such plants.

[0023] In another embodiment, a transgene introduced into a plant cell according to a method of the invention can encode a polypeptide that regulates expression from

an endogenous plant stress-regulated gene. Such a polypeptide can be, for example, a recombinantly produced polypeptide comprising a zinc finger domain, which is specific for the regulatory element, and an effector domain, which can be a repressor domain or an activator domain. The polynucleotide encoding the recombinant polypeptide can be operatively linked to and expressed from a constitutively active, inducible or tissue specific or phase specific regulatory element. Expression of the recombinant polypeptide from a plant stress-regulated promoter as disclosed herein can be particularly advantageous in that the polypeptide can be coordinately expressed with the endogenous plant stress-regulated genes upon exposure to a stress condition. The invention also provides transgenic plants produced by a method as disclosed, as well as to a plant cell obtained from such transgenic plant, wherein said plant cell exhibits altered responsiveness to the stress condition; a seed produced by the transgenic plant; and a cDNA or genomic DNA library prepared from the transgenic plant, or from a plant cell from said transgenic plant, wherein said plant cell exhibits altered responsiveness to the stress condition.

100241 In one aspect, the invention provides an isolated nucleic acid molecule comprising a nucleotide sequence substantially similar to a sequence of any one of SEQ ID NOS:2704-5379, which can alter transcription of an operatively linked polynucleotide in a plant cell in response to an abiotic stress. Additional aspects of the invention provide isolated polynucleotides, including, for example, sequences substantially similar to any of SEQ ID NOS:2704-3955, which can alter transcription of an operatively linked polynucleotide in response to a cold stress; isolated polynucleotides substantially similar to a sequence of any of SEO ID NOS:5108-5263, which can alter transcription of an operatively linked polynucleotide in response to an osmotic stress; isolated polynucleotides substantially similar to a sequence of any of SEQ ID NOS:4910-5107, which can alter transcription of an operatively linked polynucleotide in response to a saline stress; isolated polynucleotides substantially similar to a sequence of any of SEO ID NOS:4389-4654, which can alter transcription of an operatively linked polynucleotide in response to a combination of cold and osmotic stresses; isolated polynucleotides

substantially similar to a sequence of any of SEQ ID NOS:4655-4909, which can alter transcription of an operatively linked polynucleotide in response to a combination of cold and saline stresses; isolated polynucleotides substantially similar to a sequence of any of SEQ ID NOS:5264-5379, which can alter transcription of an operatively linked polynucleotide in response to a combination of osmotic and saline stresses; and isolated polynucleotides substantially similar to a sequence of any of SEQ ID NOS:3956-4388, which can alter transcription of an operatively linked polynucleotide in response to a combination of cold, osmotic and saline stresses.

[0025] Related aspects of the invention provide an isolated nucleotide sequences that can alter transcription of an operatively linked polynucleotide in response to an abiotic stress, and that hybridize under stringent conditions, preferably highly stringent conditions, to the complement of any one of SEQ ID NOS:2704-5379. Additional aspects provide an isolated nucleotide sequence that can alter transcription of an operatively linked polynucleotide in response to cold stress, and that hybridizes under stringent conditions, preferably highly stringent conditions, to the complement of any one of SEQ ID NOS:2704-3955; a nucleotide sequence that alters transcription of an operatively linked polynucleotide in response to osmotic stress, and that hybridizes under stringent conditions, preferably highly stringent conditions, to the complement of any one of SEO ID NOS:5108-5263; a nucleotide sequence that alters transcription of an operatively linked polynucleotide in response to saline stress, and that hybridizes under stringent conditions, preferably highly stringent conditions, to the complement of any one of SEQ ID NOS:4910-5107; a nucleotide sequence that alters transcription of an operatively linked polynucleotide in response to a combination of cold and osmotic stress, and that hybridizes under stringent conditions, preferably highly stringent conditions, to the complement of any one of SEQ ID NOS:4389-4654; a nucleotide sequence that alters transcription of an operatively linked polynucleotide in response to a combination of cold and saline stress, and that hybridizes under stringent conditions, preferably highly stringent conditions, to the complement of any one of SEQ ID NOS:4655-4909; a nucleotide sequence that alters transcription of an operatively linked polynucleotide in response

to an combination of osmotic and saline stress, and that hybridizes under stringent conditions, preferably highly stringent conditions, to the complement of any one of SEQ ID NOS:5264-5379; and a nucleotide sequence that alters transcription of an operatively linked polynucleotide in response to a combination of cold, osmotic and saline stress, and that hybridizes under stringent conditions, preferably highly stringent conditions, to the complement of any one of SEQ ID NOS:3956-4388.

[0026] Further aspects provide an expression cassette comprising as operatively linked components any of the above isolated nucleic acid sequences that alter transcription, a coding region, and a termination sequence. Also provided are host cells and seeds comprising such expression cassettes, plants containing such host cells and seeds and progeny of plants containing said host cells. In related aspects, the coding region of the expression cassettes comprise sequences encoding marker proteins and sequences involved in gene silencing such as antisense sequences, double stranded RNAi sequences, a triplexing agent, and sequences comprising dominant negative mutations. In additional related aspects, the coding regions comprise sequences encoding polypeptides that alter the response of a plant to an abiotic stress.

[0027] The present invention also relates to a method of modulating the responsiveness of a plant cell to a stress condition. Such a method can be performed, for example, by introducing a polynucleotide portion of a plant stress-regulated genes described herein into the plant cell, thereby modulating the responsiveness of the plant cell to a stress condition. Such a method can result in the responsiveness of the plant cell being increased upon exposure to the stress condition, which, in turn, can result in increased or decreased tolerance of the plant cell to a stress condition; or can result in the responsiveness of the plant cell to the stress condition being decreased, which, in turn, can result in increased or decreased tolerance of the plant cell to a stress condition. In one embodiment, the polynucleotide portion of the plant stress-regulated gene can integrate into the genome of the plant cell, thereby modulating the responsiveness of the plant cell to the stress condition. In another embodiment, the polynucleotide portion of the plant stress-regulated

polypeptide or functional peptide portion thereof, and can be operatively linked to a heterologous promoter. The polynucleotide portion of the plant stress-regulated gene also can contain a mutation, whereby upon integrating into the plant cell genome, the polynucleotide disrupts (knocks-out) an endogenous plant stress-regulated sequence, thereby modulating the responsiveness of the plant cell to the stress condition. Depending on whether the knocked-out gene encodes an adaptive or a maladaptive stress-regulated polypeptide, the responsiveness of the plant will be modulated accordingly.

[0028] The present invention further relates to a method of modulating the activity of a biological pathway in a plant cell, wherein the pathway involves a stressregulated polypeptide or a non-protein regulatory molecule. Such a method can be performed by introducing a polynucleotide portion of a plant stress-regulated gene, or a polynucleotide derived therefrom, for example a ribozyme derived from a nucleotide sequence as set forth in any of SEQ ID NOS:1-2703, into the plant cell, thereby modulating the activity of the biological pathway. The method can be performed with respect to a pathway involving any of the stress-regulated polypeptides as disclosed herein or encoded by the polynucleotides disclosed herein. as well as using homologs or orthologs thereof. In one embodiment, the method is performed by introducing a polynucleotide portion of a plant stress-regulated gene into the plant cell, wherein the plant stress-regulated gene comprises a nucleotide sequence as set forth in any of SEO ID NOS:1-155, 157-228, 230-232, 234-557, 559-572, 574-605, 607-634, 636-786, 788-812, 814-1262, 1264-1386, 1387-1390, 1392-1404, 1406-1444, 1446-1483, 1485-1588, 1590-1608, 1610-1633, 1634-1725, 1727-1865, 1867-1917, 1919-1927, 1929-2855, 2857-2928, 2930-2932, 2934-3256, 3258-3271, 3273-3304, 3306-3323, 3325-3333, 3335-3485, 3487-3511, 3313-3956, 3958-4078, 4080-4097, 4099-4136, 4138-4175, 4177-4279, 4281-4299, 4301-4324, 4326-4414, 4416-4552, 4554-4602, and 4604-5379, thereby modulating the activity of the biological pathway.

The present invention also relates to a method of identifying a 100291 polynucleotide that modulates a stress response in a plant cell. In one embodiment the method comprises determining gene expression in a plant exposed to at least one stress to produce an expression profile and identifying sequences whose expression is altered at least two fold compared to plants not exposed to the stress. Such an expression profile can be obtained, for example, by contacting an array of probes representative of a plant cell genome with nucleic acid molecules expressed in a plant cell exposed to the stress; and detecting one or more nucleic acid molecules expressed at a level different from a level of expression in the absence of the stress. The method can further comprise introducing the differentially expressed nucleic acid molecule into a plant cell; and detecting a modulated response of the genetically modified plant cell to a stress, thereby identifying a polynucleotide that modulates a stress response in a plant cell. The stress can be any stress, for example, an abiotic stress such as exposure to an abnormal level of cold, osmotic pressure, and salinity. The contacting is under conditions that allow for selective hybridization of a nucleic acid molecule with probe having sufficient complementarity, for example, under stringent hybridization conditions. Expression of the nucleic acid molecule can increase or decrease the tolerance of the plant cell to the stress, and the nucleic acid molecule can be expressed at a level that is less than or greater than the level of expression in the absence of the stress.

[0030] In still another embodiment, the polynucleotide portion of the plant stress-regulated gene can comprise a stress-regulated regulatory element, which can be operatively linked to a heterologous nucleotide sequence, the expression of which can modulate the responsiveness of the plant cell to a stress condition. Such a heterologous nucleotide sequence can encode, for example, a stress-inducible transcription factor such as DREB1A. The heterologous nucleotide sequence also can encode a polynucleotide that is specific for a plant stress-regulated gene, for example, an antisense molecule, an RNAi molecule, a ribozyme, and a triplexing agent, any of which, upon expression in the plant cell, reduces or inhibits expression of a stress-regulated polypeptide encoded by the gene, thereby modulating the responsiveness of

the plant cell to a stress condition, for example, an abnormal level of cold, osmotic pressure, and salinity. In another aspect, the method can include introducing a polynucleotide portion of a plant stress-regulated gene into the plant cell, wherein the plant stress-regulated gene includes a nucleotide sequence of a polynucleotide as set forth in any of SEQ ID NOS:1-155, 157-228, 230-232, 234-557, 559-572, 574-605, 607-634, 636-786, 788-812, 814-1262, 1264-1386, 1387-1390, 1392-1404, 1406-1444, 1446-1483, 1485-1588, 1590-1608, 1610-1633, 1634-1725, 1727-1865, 1867-1917, 1919-1927, 1929-2855, 2857-2928, 2930-2932, 2934-3256, 3258-3271, 3273-3304, 3306-3323, 3325-3333, 3335-3485, 3487-3511, 3313-3956, 3958-4078, 4080-4097, 4099-4136, 4138-4175, 4177-4279, 4281-4299, 4301-4324, 4326-4414, 4416-4552, 4554-4602, and 4604-5379, thereby modulating the responsiveness of the plant cell to a stress condition. The invention also relates to a plant cell obtained by any of the methods of modulating the responsiveness of a plant to a stress condition or combination of stress conditions, and to a plant comprising such a plant cell.

The present invention further relates to a method of selecting a plant having [0031] an altered resistance to an abiotic stress condition or a combination of abiotic stress conditions, such a method being useful for marker-assisted breeding. Such a method can be performed, for example, by contacting nucleic acid molecules representative of expressed polynucleotides in a plant cell of a plant to be examined for having an altered resistance to an abiotic stress with a nucleic acid probes that selectively hybridizes under stringent conditions to a plant stress-regulated gene comprising a nucleotide sequence as set forth in any of SEQ ID NO:1-5379; detecting a level of selective hybridization of the nucleic acid probes to a nucleic acid molecule representative of an expressed polynucleotide in the plant cell, wherein the level of selective hybridization corresponds to the level of the expressed polynucleotide in the plant cell, which is indicative of resistance of the plant to an abiotic stress; and selecting a plant having a level of expression of a polynucleotide indicative of altered resistance to an abiotic stress condition. For example, the abiotic stress condition can be cold stress, and the nucleic acid probe can include at least about 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID NOS:1-1261 and 2704-3955, for

example, at least about 15 nucleotides of a nucleotide sequence as set forth in any of SEO ID NOS:1-155, 157-228, 230-232, 234-557, 559-572, 574-605, 607-634, 636-786, 788-812, 814-1261, 2704-2855, 2857-2928, 2930-2932, 2934-3256, 3258-3271, 3273-3304, 3306-3323, 3325-3333, 3335-3485, 3487-3511. and 3313-3955; or the abiotic stress condition can be saline stress, and the nucleic acid probe can include at least about 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID NOS:2226-2427 and 4910-5107; or the abiotic stress condition can be osmotic stress, and the nucleic acid probe can include at least about 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID NOS:2428-2585 and 5108-5263. In addition, a combination of abiotic stress conditions can be a combination of cold stress and osmotic stress, and the nucleic acid probe can include at least about 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID NOS:1669-1969 and 4389-4654, for example, at least about 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID NOS:1699-1725, 1727-1865, 1867-1917, 1919-1927, 1929-1969, 4389-4414, 4416-4552, 4554-4602, 4604-4612, and 4613-4654; or the combination of abiotic stress conditions can be a combination of cold stress and saline stress, and the nucleic acid probe can include at least about 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID NOS:1970-2226 and 4655-4909; or the combination of abiotic stress conditions can be a combination of osmotic stress and saline stress, and the nucleic acid probe can include at least about 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID NOS:2586-2703 and 5264-5379; or the combination of abiotic stress conditions can be a combination of cold stress, osmotic stress and saline stress, and the nucleic acid probe can include at least about 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID NOS:1262-1698 and 3956-4388, for example, at least about 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID NOS:1262, 1264-1386, 1387-1390, 1392-1404, 1406-1444, 1446-1483, 1485-1588, 1590-1608, 1610-1633, 1634-1698, 3956, 3958-4078, 4080-4097, 4099-4136, 4138-4175, 4177-4279,4281-4299, 4301-4324, and 4326-4388.

The present invention also relates to a method of expressing a heterologous 100321 nucleotide sequence in a plant cell. Such a method can be performed, for example, by introducing into the plant cell a plant stress-regulated regulatory element operatively linked to the heterologous nucleotide sequence, whereby, upon exposure of the plant cell to a stress condition, the heterologous nucleotide sequence is expressed in the plant cell. In one embodiment, the stress-regulated gene regulatory element is any of the sequences described herein that are capable of altering transcription of an operatively linked sequence in response to an abiotic stress, for example, SEQ ID NOS:2704-5379. In another embodiment, stress-regulated gene regulatory element comprises a nucleotide sequence as set forth in any of SEQ ID NOS:2704-2855, 2857-2928, 2930-2932, 2934-3256, 3258-3271, 3273-3304, 3306-3323, 3325-3333, 3335-3485, 3487-3511, 3313-3956, 3958-4078, 4080-4097, 4099-4136, 4138-4175,4177-4279, 4281-4299, 4301-4324, 4326-4414, 4416-4552, 4554-4602, and 4604-5379, whereby, upon exposure of the plant cell to stress condition, the heterologous nucleotide sequence is expressed in the plant cell. The heterologous nucleotide sequence can encode a selectable marker, a diagnostic marker, or a polypeptide that confers a desirable trait upon the plant cell, for example, a polypeptide that improves the nutritional value, digestibility or ornamental value of the plant cell, or a plant comprising the plant cell.

[0033] The present invention additionally relates to a method of identifying a stress condition to which a plant cell was exposed by comparing an expression profile from a test plant suspected of having been exposed to at least one stress condition to an expression profile obtained from a reference plant, preferably of the same species, which has been exposed to the suspected stress condition. Such a method can be performed, for example, by contacting nucleic acid molecules representative of expressed polynucleotides in cells of the test plant with at least one nucleic acid probe under conditions suitable for selective hybridization to a complementary nucleotide sequence, wherein the probe comprises at least 15 nucleotides of a plant stress-regulated gene, wherein the stress-regulated gene does not have a nucleotide sequence of a polynucleotide as set forth in any of SEQ ID NOS:156, 229, 233, 558, 573, 606,

635, 787, 813, 1263, 1386, 1391, 1405, 1445, 1484, 1589, 1609, 1634, 1726, 1866, 1918 or 1928, or a nucleotide sequence complementary thereto, whereby detecting selective hybridization of at least one nucleic acid probe, or detecting a change in a level of selective hybridization as compared to a level of selective hybridization obtained using nucleic acid molecules representative of expressed polynucleotides in cells of a plant known not have been exposed to an abiotic stress, indicates that the test plant has been exposed to an abiotic stress, and whereby an absence of selective hybridization of at least one nucleic acid probe indicates that the test plant has not been exposed to an abiotic stress. For example, the abiotic stress is cold stress, and the probe can include at least 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID NOS:1-155, 157-228, 230-232, 234-557, 559-572, 574-605, 607-634, 636-786, 788-812, 814-1261 or a nucleotide sequence complementary thereto; or the abiotic stress can be a saline stress, and the probe can include at least 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID NOS:2226-2427 or a nucleotide sequence complementary thereto; or the abiotic stress can be an osmotic stress, and the probe can include at least 15 nucleotides of a nucleotide sequence as set forth in two or more of SEQ ID NOS:2428-2585 or a nucleotide sequence complementary thereto.

[0034] A method of identifying a stress condition to which a plant cell was exposed also can be performed, for example, by contacting nucleic acid molecules expressed in the test plant cell with an array of probes representative of the plant cell genome; detecting a profile of expressed nucleic acid molecules characteristic of a stress response, and comparing the expression pattern in the test plant to the expression pattern obtained from a reference plant thereby identifying the stress condition to which the plant cell was exposed. The contacting is under conditions that allow for selective hybridization of a nucleic acid molecule with probes having sufficient complementarity, for example, under stringent hybridization conditions. The profile can be characteristic of exposure to a single stress condition, for example, an abnormal level of cold, osmotic pressure, or salinity, or can be characteristic of exposure to more than one stress condition, for example, cold, increased osmotic

pressure and increased salinity. In one embodiment, the nucleotide sequence of a gene whose expression is detected is selected from a polynucleotide comprising any of SEQ ID NOS:1-2703. In further embodiments, the nucleotide sequence of a gene that is expressed in response a particular stress or combination of stresses can comprise a polynucleotide expressed in response to cold stress (SEQ ID NOS:1-1261), osmotic stress (SEQ ID NOS:2428-2585), saline (salt) stress (SEQ ID NOS:2227-2427), a combination of cold and osmotic stress (SEQ ID NOS:1699-1969), a combination of saline and osmotic stress (SEQ ID NOS:1970-2226), a combination of osmotic and saline stress (SEQ ID NOS:2586-2703), or a combination of cold, osmotic and saline stress (SEQ ID NOS:1262-1698).

In another embodiment, the method can be used for determining whether a 100351 test plant has been exposed to a combination of abiotic stress conditions. Such a method can be performed, for example, by contacting nucleic acid molecules representative of expressed polynucleotides in cells of the test plant with at least one nucleic acid probe under conditions suitable for selective hybridization to a complementary nucleotide sequence, whereby detecting selective hybridization of at least one nucleic acid probe, or detecting a change in a level of selective hybridization as compared to a level of selective hybridization obtained using nucleic acid molecules representative of expressed polynucleotides in cells of a plant known not have been exposed to a combination of stress conditions, indicates that the test plant has been exposed to a combination of abiotic stress conditions, and whereby an absence of selective hybridization of at least one nucleic acid probe indicates that the test plant has not been exposed to a combination of abiotic stress conditions. For example, the combination of abiotic stress conditions can be a combination of a cold stress and an osmotic stress, and the probe can include at least 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID NOS:1699-1969, or a nucleotide sequence complementary thereto; or the combination of abiotic stress conditions can be a combination of a cold stress and a saline stress, and the probe can include at least 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID

NOS:1970-2226, or a nucleotide sequence complementary thereto; or the combination of abiotic stress conditions can be a combination of an osmotic stress and a saline stress, and the probe can included at least 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID NOS:2586-2703, or a nucleotide sequence complementary thereto; or the combination of abiotic stress conditions can be a combination of a cold stress, a saline stress and an osmotic stress, and the probe can include at least 15 nucleotides of a nucleotide sequence as set forth in any of SEQ ID NOS:1262-1698, or a nucleotide sequence complementary thereto.

[0036] The present invention also relates to a method for monitoring a population of plants for exposure to a stress condition or combination of stress conditions. Such a method can be performed, for example, by introducing into the population of a plants a sentinel plant, wherein said sentinel plant is a transgenic plant, which contains plant cells containing a stress-regulated regulatory element operatively linked to a polynucleotide encoding a detectable marker; and examining the sentinel plant for expression of the detectable marker, which is indicative of exposure of the population of plants to a stress condition or combination of stress conditions. The stress condition or combination of stress conditions can be any such condition or conditions, particularly an abiotic stress condition or combination of abiotic stress conditions. The detectable marker can be any reporter molecule that is readily or conveniently detectable, particularly a marker that is visibly detectable, for example, a luminescent detectable marker such as luciferin, or a fluorescent detectable marker such as a green fluorescent protein, a yellow fluorescent protein, a cyan fluorescent protein, a red fluorescent protein, or an enhanced or modified form thereof.

[0037] The present invention further relates to a transgenic plant, which contains a nucleic acid construct comprising a polynucleotide portion of plant stress-regulated polynucleotide. In one embodiment, the transgenic plant exhibits altered responsiveness to a stress condition as compared to a corresponding reference plant not containing the construct. Such a transgenic plant can contain, for example, a construct that disrupts an endogenous stress-regulated gene in the plant, thereby

reducing or inhibiting expression of the gene in response to a stress condition. Such a knock-out can increase or decrease tolerance of the plant to a stress condition. The transgene also can comprise a coding sequence of a plant stress-regulated gene, which can be operatively linked to a heterologous regulatory element such as a constitutively active regulatory element, an regulated regulatory element, a tissues specific or phase specific regulatory element, or the like. In another embodiment, the transgenic plant contains a nucleic acid construct comprising a plant stress-regulated regulatory element, which can be operatively linked to a heterologous nucleotide sequence that can encode a polypeptide. Expression of the heterologous polypeptide can confer a desirable characteristic on the plant, for example, can improve the nutritional or ornamental value of the transgenic plant. In still another embodiment, the transgenic plant contains multiple nucleic acid constructs, which can be multiple copies of the same construct, or can be two or more different constructs.

The present invention also relates to a plant stress-regulated regulatory [0038] element, which is obtained from a plant stress-regulated polynucleotide disclosed herein for example any of SEQ ID NOS:2704-5379; a homolog or ortholog thereof. The invention also provides a method of identifying an agent, for example a transcription factor, that specifically binds to or activates a plant stress-regulated regulatory element. Such a method can be performed, for example, by contacting the regulatory element with a plant cell extract, and identifying polypeptides that specifically bind to the regulatory element. Confirmation that the specifically binding polypeptide is a transcription factor can be demonstrated using, for example, the stress-regulated regulatory element operably linked to a reporter gene, and detecting expression of the reporter gene. Control constructs comprising a regulatory element, other than a plant stress-regulated regulatory element, operatively linked to a reporter molecule can be used to confirm that the transcription factor is specific for the plant stress-regulated regulatory element. A polynucleotide encoding such a transcription factor also can be obtained.

The present invention also relates to a method of using a polynucleotide 100391 portion of a plant stress-regulated gene to confer a selective advantage on a plant cell. In one embodiment, such a method is performed by introducing a plant stressregulated regulatory element into a plant cell such as those described herein, wherein, upon exposure of the plant cell to a stress condition to which the regulatory element is responsive, a nucleotide sequence operatively linked to the regulatory element is expressed, thereby conferring a selective advantage to plant cell. The operatively linked nucleotide sequence can be, for example, a transcription factor, the expression of which induces the further expression of polynucleotides involved in a stress response, thereby enhancing the response of a plant to the stress condition. In another embodiment, a coding sequence of a plant stress-regulated gene as disclosed herein is introduced into the cell, thereby providing the plant with a selective advantage in response to a stress condition. In still another embodiment, the method results in the knock-out of a plant stress-regulated gene as disclosed herein in a first population of plants, thereby providing a selective advantage to a stress condition in a second population of plants.

[0040] The invention further relates to a method of identifying an agent that modulates the activity of a stress-regulated regulatory element of a plant. In a particular embodiment, is provided a method for identifying an agent that alters the activity of an abiotic stress responsive regulatory element comprising contacting the agent or a composition containing an agent to be tested with at least one abiotic stress responsive regulatory element, preferably selected from the group consisting of SEQ ID NOS:2704-5379 (see Table 2), and determining the effect of the agent on the ability of the regulatory sequence to regulate transcription. In further embodiments, the regulatory elements are associated with particular stresses or combination of stresses such as cold stress (SEQ ID NOS:2704-3955), osmotic stress (SEQ ID NOS:5108-5263), saline stress (SEQ ID NOS:4910-5107), a combination of cold and osmotic stress (SEQ ID NOS:4389-4654), a combination of cold and saline stress (SEQ ID NOS:4655-4909), a combination of osmotic and saline stress (SEQ ID NOS:5264-5379), or a combination of cold, osmotic and saline stress (SEQ ID NOS:5264-5379), or a combination of cold, osmotic and saline stress (SEQ ID

NOS:3956-4388). In one embodiment, the regulatory element can be operatively linked to a heterologous polynucleotide encoding a reporter molecule, and an agent that modulates the activity of the stress-regulated regulatory element can be identified by detecting a change in expression of the reporter molecule due to contacting the regulatory element with the agent. Such a method can be performed *in vitro* in a plant cell-free system, or in a plant cell in culture or in a plant in situ. In another embodiment, the agent is contacted with a transgenic plant containing an introduced plant stress-regulated regulatory element, and an agent that modulates the activity of the regulatory element is identified by detecting a phenotypic change in the transgenic plant. The methods of the invention can be performed in the presence or absence of the stress condition to which the particularly regulatory element is responsive.

Another aspect provides a method for identifying an agent that alters [0041] abiotic stress responsive polynucleotide expression in a plant or plant cell comprising contacting a plant or plant cell with a test agent; subjecting the plant cell or plant cell to an abiotic stress or combination of stresses before, during or after contact with the agent to be tested; obtaining an expression profile of the plant or plant cell and comparing the expression profile of the plant or plant cell to an expression profile from a plant or plant cell not exposed to the abiotic stress or combination of stresses. In one embodiment, the expression profile comprises expression data for at least one nucleotide sequence comprising any of SEQ ID NOS:1-5379 (see Tables 1 and 2). In additional embodiments, the expression profile comprises expression data for at least one, and preferably two or more sequences associated with a particular abiotic stress or combination of stresses such as cold stress (SEQ ID NOS:1-1261 and 2704-3955), osmotic stress (SEQ ID NOS:2428-2585 and 5108-5263), saline stress (SEQ ID NOS:2227-2427 and 4910-5107), a combination of cold and osmotic stress (SEQ ID NOS:1699-1969 and 4389-4654), a combination of cold and saline stress (SEQ ID NOS:1970-2226 and 4655-4909), a combination of osmotic and saline stress (SEQ ID NOS:2586-2703 and 5264-5379), or a combination of cold, osmotic and saline stress (SEO ID NOS:1262-1698 and 3956-4388).

[0042] Still another aspect provides nucleotide probes useful for detecting an abiotic stress response in plants, the probes comprising a nucleotide sequence of at least 15, 25, 50 or 100 nucleotides that hybridizes under stringent, preferably highly stringent, conditions to at least one sequence comprising any of SEQ ID NOS:1-2703. Also provided are nucleotide probes comprising at least 15, 25, 50 or 100 nucleotides in length that hybridize under stringent, preferably highly stringent conditions, to at least one gene associated with a particular stress or combination of stresses, for example cold stress, (SEQ ID NOS:1-1261), osmotic stress (SEQ ID NOS:2428-2585), saline stress (SEQ ID NOS:2227-2427), a combination of cold and osmotic stress (SEQ ID NOS:1699-1969), a combination of cold and saline stress (SEQ ID NOS:2586-2703), or a combination of cold, osmotic, and saline stress (SEQ ID NOS:1262-1698).

An additional aspect provides a method for marker-assisted breeding to [0043] select plants having an altered resistance to abiotic stress comprising obtaining nucleic acid molecules from the plants to be selected; contacting the nucleic acid molecules with one or more probes that selectively hybridize under stringent, preferably highly stringent, conditions to a nucleic acid sequence selected from the group consisting of SEO ID NOS:1-2703; detecting the hybridization of the one or more probes to the nucleic acid sequences wherein the presence of the hybridization indicates the presence of a gene associated with altered resistance to abiotic stress; and selecting plants on the basis of the presence or absence of such hybridization. Marker-assisted selection can also be accomplished using one or more probes which selectively hybridize under stringent, preferably highly stringent conditions, to a nucleotide sequence comprising a polynucleotide expressed in response associated with a particular stress, for example, a nucleotide sequence comprising any of SEQ ID NOS:1-1261 (cold stress), SEQ ID NOS:2428-2585 (osmotic stress), SEQ ID NOS:2227-2427 (saline stress), SEQ ID NOS:1699-1969 (cold and osmotic stress), SEO ID NOS:1970-2226 (cold and saline stress), SEQ ID NOS:2586-2703 (osmotic and saline stress), or SEQ ID NOS:1262-1698 (cold, osmotic and saline stress). In

each case marker-assisted selection can be accomplished using a probe or probes to a single sequence or multiple sequences. If multiple sequences are used they can be used simultaneously or sequentially.

[0044] A further aspect provides a method for monitoring a population of plants comprising providing at least one sentinel plant containing a recombinant polynucleotide comprising a stress responsive regulatory sequence selected from the group consisting of SEQ ID NOS:2704-5379 which is operatively linked to a nucleotide sequence encoding a detectable marker, for example a fluorescent protein. Additional aspects provide the use of various regulatory sequences including those associated with cold stress (SEQ ID NOS:2704-3955), osmotic stress (SEQ ID NOS:5108-5263), saline stress (SEQ ID NOS:4910-5107), cold and osmotic stress (SEQ ID NOS:4389-4654), cold and saline stress (SEQ ID NOS:4655-4909), osmotic and saline stress (SEQ ID NOS:3956-4388), or fragments thereof wherein such fragments can alter transcription of an operatively linked nucleotide sequence in response to an abiotic stress.

[0045] A further aspect provides a computer readable medium having stored thereon computer executable instructions for performing a method comprising receiving data on gene expression in a test plant of at least one nucleic acid molecule having at least 70%, preferably at least 80%, more preferably at least 90%, and most preferably at least 95% nucleotide sequence identity to one or more polynucleotide sequences as set forth in any of SEQ ID NOS:1-2703; and comparing expression data from the test plant to expression data for the same polynucleotide sequence or sequences in a plant that has been exposed to at least one abiotic stress.

[0046] Yet a further aspect provides a computer readable medium having stored thereon a data structure comprising, sequence data for at least one, and preferably a plurality of nucleic acid molecules having at least 70%, preferably at least 80%, more preferably at least 90%, and most preferably at least 95% nucleotide sequence identity

to a polynucleotide comprising any of SEQ ID NOS:1-2703, or the complement thereof; and a module receiving the nucleic acid molecule sequence data which compares the nucleic acid molecule sequence data to at least one other nucleic acid sequence.

DETAILED DESCRIPTION OF THE INVENTION

[0047] The present invention relates to clusters of genes that are induced in response to one or a combination of abiotic stress conditions. Abiotic stress conditions, such as a shortage or excess of solar energy, water and nutrients, and salinity, high and low temperature, or pollution (e.g., heavy metals), can have a major impact on plant growth and can significantly reduce the yield, for example, of cultivars. Under conditions of abiotic stress, the growth of plant cells is inhibited by arresting the cell cycle in late G1, before DNA synthesis, or at the G2/M boundary (see Dudits, Plant Cell Division, Portland Press Research, Monograph; Francis, Dudits, and Inze, eds., 1997; chap. 2, page 21; Bergounioux, <u>Protoplasma</u>
142:127-136, 1988). The identification of stress-regulated gene clusters, using microarray technology, provides a means to identify plant stress-regulated genes.

[0048] As used herein, the term "cluster," when used in reference to stress-regulated genes, refers to nucleotide sequences of genes that have been selected by drawing Venn diagrams, and selecting those genes that are regulated only by a selected stress condition. In general, a cluster of stress-regulated genes includes at least 5, 10, 15, or 20 genes, including polynucleotide portions thereof, each of which is responsive to the same selected stress condition or conditions. The selected stress condition can be a single stress condition, for example, cold, osmotic stress or salinity stress (see Tables 3-14), or can be a selected combination of stress conditions, for example, cold, osmotic stress and salinity stress (see Tables 15-26). In addition, a cluster can be selected based on specifying that all of the genes are coordinately regulated, for example, they all start at a low level and are induced to a higher level. However, a cluster of saline stress-regulated genes, for example, that was selected for coordinate regulation from low to high, also can be decreased in response to cold or

mannitol. By varying the parameters used for selecting a cluster of gene nucleotide sequences, those genes that are expressed in a specific manner following a stress can be identified.

[0049] As used herein in reference to a polynucleotide or polynucleotide portion of a gene or nucleic acid molecule, the term "isolated" means a polynucleotide, polynucleotide portion of a gene, or nucleic acid molecule that is free of one or both of the nucleotide sequences that normally flank the polynucleotide in a genome of a naturally-occurring organism from which the polynucleotide is derived. The term includes, for example, a polynucleotide or fragment thereof that is incorporated into a vector or expression cassette; into an autonomously replicating plasmid or virus; into the genomic DNA of a prokaryote or eukaryote; or that exists as a separate molecule independent of other polynucleotides. It also includes a recombinant polynucleotide that is part of a hybrid polynucleotide, for example, one encoding a polypeptide sequence.

[0050] The terms "polynucleotide," "oligonucleotide," and "nucleic acid sequence" are used interchangeably herein to refer to a polymeric (2 or more monomers) form of nucleotides of any length, either ribonucleotides or deoxyribonucleotides. Although nucleotides are usually joined by phosphodiester linkages, the term also includes polymers containing neutral amide backbone linkages composed of aminoethyl glycine units. The terms are used only to refer to the primary structure of the molecule. Thus, the term includes double stranded and single stranded DNA molecules, including a sense strand or an antisense strand, and RNA molecules as well as genomic DNA, cDNA, mRNA and the like. It will be recognized that such polynucleotides can be modified, for example, by including a label such as a radioactive, fluorescent or other tag, by methylation, by the inclusion of a cap structure, by containing a substitution of one or more of the naturally occurring nucleotides with a nucleotide analog, by containing an internucleotide modification such as having uncharged linkages (e.g., methyl phosphonates, phosphotriesters, phosphoramidates, carbamates, or the like), by containing a pendant moiety such as a

protein (e.g., a nuclease, toxin, antibody, signal peptide, poly-L-lysine, or the like), by containing an intercalator such as acridine or psoralen, by containing a chelator, which can be a metal such as boron, an oxidative metal, or a radioactive metal, by containing an alkylator, or by having a modified linkage (e.g., an alpha anomeric nucleic acid).

[0051] The term "recombinant nucleic acid molecule" refers to a polynucleotide produced by human intervention. A recombinant nucleic acid molecule can contain two or more nucleotide sequences that are linked in a manner such that the product is not found in a cell in nature. In particular, the two or more nucleotide sequences can be operatively linked and, for example, can encode a fusion polypeptide, or can comprise a nucleotide sequence and a regulatory element. A recombinant nucleic acid molecule also can be based on, but different, from a naturally occurring polynucleotide, for example, a polynucleotide having one or more nucleotide changes such that a first codon, which normally is found in the polynucleotide, is replaced with a degenerate codon that encodes the same or a conservative amino acid, or such that a sequence of interest is introduced into the polynucleotide, for example, a restriction endonuclease recognition site or a splice site, a promoter, a DNA replication initiation site, or the like.

[0052] As used herein, the term "abiotic stress" or "abiotic stress condition" refers to the exposure of a plant, plant cell, or the like, to a non-living ("abiotic") physical or chemical agent or condition that has an adverse effect on metabolism, growth, development, propagation and/or survival of the plant (collectively "growth"). An abiotic stress can be imposed on a plant due, for example, to an environmental factor such as water (e.g., flooding, drought, dehydration), anaerobic conditions (e.g., a low level of oxygen), abnormal osmotic conditions, salinity or temperature (e.g., hot/heat, cold, freezing, frost), a deficiency of nutrients or exposure to pollutants, or by a hormone, second messenger or other molecule. Anaerobic stress, for example, is due to a reduction in oxygen levels (hypoxia or anoxia) sufficient to produce a stress response. A flooding stress can be due to prolonged or transient immersion of a plant,

plant part, tissue or isolated cell in a liquid medium such as occurs during monsoon, wet season, flash flooding or excessive irrigation of plants, or the like. A cold stress or heat stress can occur due to a decrease or increase, respectively, in the temperature from the optimum range of growth temperatures for a particular plant species. Such optimum growth temperature ranges are readily determined or known to those skilled in the art. Dehydration stress can be induced by the loss of water, reduced turgor, or reduced water content of a cell, tissue, organ or whole plant. Drought stress can be induced by or associated with the deprivation of water or reduced supply of water to a cell, tissue, organ or organism. Saline stress (salt stress) can be associated with or induced by a perturbation in the osmotic potential of the intracellular or extracellular environment of a cell. Osmotic stress also can be associated with or induced by a change, for example, in the concentration of molecules in the intracellular or extracellular environment of a plant cell, particularly where the molecules cannot be partitioned across the plant cell membrane.

[0053] As disclosed herein, clusters of plant stress-regulated genes (Example 1; see, also, Tables 1-31) and homologs and orthologs thereof (Table 32) have been identified. Remarkably, several of the stress-regulated genes previously were known to encode polypeptides having defined cellular functions, including roles as transcription factors, enzymes such as kinases, and structural proteins such as channel proteins (see Tables 29-31). The identification of Arabidopsis stress-regulated genes provides a means to identify homologous and orthologous genes and gene sequences in other plant species using well known procedures and algorithms based on identity (or homology) to the disclosed sequences. Thus, the invention provides polynucleotide sequences comprising plant stress-regulated genes that are homologs or orthologs, variants, or otherwise substantially similar to the polynucleotides disclosed herein, and having an E value $\leq 1 \times 10^{-8}$, which can be identified, for example, by a BLASTN search using the Arabidopsis polynucleotides of Tables 1 and 2 (SEO ID NOS:1-5379) as query sequences (see Table 32, on CD).

[0054] A polynucleotide sequence of a stress-regulated gene as disclosed herein can be particularly useful for performing the methods of the invention on a variety of plants, including but not limited to, corn (Zea mays), Brassica sp. (e.g., B. napus, B. rapa, B. juncea), particularly those Brassica species useful as sources of seed oil, alfalfa (Medicago sativa), rice (Oryza sativa), rye (Secale cereale), sorghum (Sorghum bicolor, Sorghum vulgare), millet (e.g., pearl millet (Pennisetum glaucum), proso millet (Panicum miliaceum), foxtail millet (Setaria italica), finger millet (Eleusine coracana)), sunflower (Helianthus annuus), safflower (Carthamus tinctorius), wheat (Triticum aestivum), soybean (Glycine max), tobacco (Nicotiana tabacum), potato (Solanum tuberosum), peanuts (Arachis hypogaea), cotton (Gossypium barbadense, Gossypium hirsutum), sweet potato (Ipomoea batatus), cassava (Manihot esculenta), coffee (Cofea spp.), coconut (Cocos nucifera), pineapple (Ananas comosus), citrus trees (Citrus spp.), cocoa (Theobroma cacao), tea (Camellia sinensis), banana (Musa spp.), avocado (Persea ultilane), fig (Ficus casica), guava (Psidium guajava), mango (Mangifera indica), olive (Olea europaea), papaya (Carica papaya), cashew (Anacardium occidentale), macadamia (Macadamia integrifolia), almond (Prunus amvgdalus), sugar beets (Beta vulgaris), sugarcane (Saccharum spp.), oats, duckweed (Lemna), barley, tomatoes (Lycopersicon esculentum), lettuce (e.g., Lactuca sativa), green beans (Phaseolus vulgaris), lima beans (Phaseolus limensis), peas (Lathyrus spp.), and members of the genus Cucumis such as cucumber (C. sativus), cantaloupe (C. cantalupensis), and musk melon (C. melo), Ornamentals such as azalea (Rhododendron spp.), hydrangea (Macrophylla hydrangea), hibiscus (Hibiscus rosasanensis), roses (Rosa spp.), tulips (Tulipa spp.), daffodils (Narcissus spp.), petunias (Petunia hybrida), carnation (Dianthus caryophyllus), poinsettia (Euphorbia pulcherrima), and chrysanthemum are also included. Additional ornamentals within the scope of the invention include impatiens, Begonia, Pelargonium, Viola, Cyclamen, Verbena, Vinca, Tagetes, Primula, Saint Paulia, Agertum, Amaranthus, Antihirrhinum, Aquilegia, Cineraria, Clover, Cosmo, Cowpea, Dahlia, Datura, Delphinium, Gerbera, Gladiolus, Gloxinia, Hippeastrum, Mesembryanthemum, Salpiglossos, and Zinnia. Conifers that may be employed in

practicing the present invention include, for example, pines such as loblolly pine (Pinus taeda), slash pine (Pinus elliotii), ponderosa pine (Pinus ponderosa), lodgepole pine (Pinus contorta), and Monterey pine (Pinus radiata), Douglas-fir (Pseudotsuga menziesii); Western hemlock (Tsuga ultilane); Sitka spruce (Picea glauca); redwood (Sequoia sempervirens); true firs such as silver fir (Abies amabilis) and balsam fir (Abies balsamea); and cedars such as Western red cedar (Thuja plicata) and Alaska yellow-cedar (Chamaecyparis nootkatensis).

[0055] Leguminous plants which may be used in the practice of the present invention include beans and peas. Beans include guar, locust bean, fenugreek, soybean, garden beans, cowpea, mung bean, lima bean, fava bean, lentils, chickpea, etc. Legumes include, but are not limited to, Arachis, e.g., peanuts, Vicia, e.g., crown vetch, hairy vetch, adzuki bean, mung bean, and chickpea, Lupinus, e.g., lupine, trifolium, Phaseolus, e.g., common bean and lima bean, Pisum, e.g., field bean, Melilotus, e.g., clover, Medicago, e.g., alfalfa, Lotus, e.g., trefoil, lens, e.g., lentil, and false indigo. Preferred forage and turf grass for use in the methods of the invention include alfalfa, orchard grass, tall fescue, perennial ryegrass, creeping bent grass, and redtop.

[0056] Other plants within the scope of the invention include Acacia, aneth, artichoke, arugula, blackberry, canola, cilantro, clementines, escarole, eucalyptus, fennel, grapefruit, honey dew, jicama, kiwifruit, lemon, lime, mushroom, nut, okra, orange, parsley, persimmon, plantain, pomegranate, poplar, radiata pine, radicchio, Southern pine, sweetgum, tangerine, triticale, vine, yams, apple, pear, quince, cherry, apricot, melon, hemp, buckwheat, grape, raspberry, chenopodium, blueberry, nectarine, peach, plum, strawberry, watermelon, eggplant, pepper, cauliflower, Brassica, e.g., broccoli, cabbage, ultilan sprouts, onion, carrot, leek, beet, broad bean, celery, radish, pumpkin, endive, gourd, garlic, snapbean, spinach, squash, turnip, ultilane, chicory, groundnut and zucchini.

As used herein, the term "substantially similar", when used herein with respect to a nucleotide sequence, means a nucleotide sequence corresponding to a reference nucleotide sequence, wherein the corresponding sequence encodes a polypeptide or comprises a regulatory element having substantially the same structure and function as the polypeptide encoded by the reference nucleotide sequence, for example, where only changes in amino acids not affecting the polypeptide function occur. For purposes of the present invention, a reference (or query) sequence is a polynucleotide sequence as set forth in any of SEQ ID NOS:1-2703 or a polyneptide encoded thereby. Desirably, a substantially similar nucleotide sequence encodes the polypeptide encoded by the reference nucleotide sequence. The percentage of identity between the substantially similar nucleotide sequence and the reference nucleotide sequence desirably is at least 60%, more desirably at least 75%, preferably at least 90%, more preferably at least 95%, still more preferably at least 99% and including 100%. A nucleotide sequence is "substantially similar" to reference nucleotide sequence hybridizes to the reference nucleotide sequence in 7% sodium dodecyl sulfate (SDS), 0.5 M NaPO4, 1 mM EDTA at 50°C with washing in 2X SSC, 0.1% SDS at 50°C, more desirably in 7% sodium dodecyl sulfate (SDS), 0.5 M NaPO₄, 1 mM EDTA at 50°C with washing in 1X SSC, 0.1% SDS at 50°C (stringent conditions), more desirably still in 7% sodium dodecyl sulfate (SDS). 0.5 M NaPO₄, 1 mM EDTA at 50°C with washing in 0.5X SSC, 0.1% SDS at 50°C (high stringency), preferably in 7% sodium dodecyl sulfate (SDS), 0.5 M NaPO₄, 1 mM EDTA at 50°C with washing in 0.1X SSC, 0.1% SDS at 50°C (very high stringency), more preferably in 7% sodium dodecyl sulfate (SDS), 0.5 M NaPO₄, 1 mM EDTA at 50°C with washing in 0.1X SSC, 0.1% SDS at 65°C (extremely high stringency).

[0058] In addition, the term "substantially similar," when used in reference to a polypeptide sequence, means that an amino acid sequence relative to a reference (query) sequence shares at least about 65% amino acid sequence identity, particularly at least about 75% amino acid sequence identity, and preferably at least about 85%,

more preferably at least about 90%, and most preferably at least about 95% or greater amino acid sequence identity. Generally, sequences having an $E \le 10^{-8}$ are considered to be substantially similar to a query sequence. Such sequence identity can take into account conservative amino acid changes that do not substantially affect the function of a polypeptide. As such, homologs or orthologs of the *Arabidopsis* stress-regulated nucleotide sequences disclosed herein, variants thereof, and polypeptides substantially similar to the polynucleotide sequence of *Arabidopsis* stress-regulated genes set forth in SEQ ID NOS:1-5379 are encompassed within the present invention and, therefore, useful for practicing the methods of the invention (see, for example, Table 32, which is on the CD-R filed herewith, and incorporated herein by reference).

[0059] Homology or identity is often measured using sequence analysis software such as the Sequence Analysis Software Package of the Genetics Computer Group (University of Wisconsin Biotechnology Center, 1710 University Avenue, Madison, WI 53705). Such software matches similar sequences by assigning degrees of homology to various deletions, substitutions and other modifications. The terms "homology" and "identity," when used herein in the context of two or more nucleic acids or polypeptide sequences, refer to two or more sequences or subsequences that are the same or have a specified percentage of amino acid residues or of nucleotides that are the same when compared and aligned for maximum correspondence over a comparison window or designated region as measured using any number of sequence comparison algorithms or by manual alignment and visual inspection.

[0060] For sequence comparison, typically one sequence acts as a reference sequence, to which test sequences are compared. When using a sequence comparison algorithm, test and reference sequences are entered into a computer, subsequence coordinates are designated, if necessary, and sequence algorithm program parameters are designated. Default program parameters can be used, or alternative parameters can be designated. The sequence comparison algorithm then calculates the percent sequence identities for the test sequences relative to the reference sequence, based on the program parameters.

The term "comparison window" is used broadly herein to include reference to a segment of any one of the number of contiguous positions, for example, about 20 to 600 positions, for example, amino acid or nucleotide position, usually about 50 to about 200 positions, more usually about 100 to about 150 positions, in which a sequence may be compared to a reference sequence of the same number of contiguous positions after the two sequences are optimally aligned. Methods of alignment of sequence for comparison are well-known in the art. Optimal alignment of sequences for comparison can be conducted, for example, by the local homology algorithm of Smith and Waterman (Adv. Appl. Math. 2:482, 1981), by the homology alignment algorithm of Needleman and Wunsch (J. Mol. Biol. 48:443, 1970), by the search for similarity method of Person and Lipman (Proc. Natl. Acad. Sci., USA 85:2444, 1988), each of which is incorporated herein by reference; by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Dr., Madison, WI); or by manual alignment and visual inspection. Other algorithms for determining homology or identity include, for example, in addition to a BLAST program (Basic Local Alignment Search Tool at the National Center for Biological Information), ALIGN, AMAS (Analysis of Multiply Aligned Sequences), AMPS (Protein Multiple Sequence Alignment), ASSET (Aligned Segment Statistical Evaluation Tool), BANDS, BESTSCOR, BIOSCAN (Biological Sequence Comparative Analysis Node), BLIMPS (BLocks IMProved Searcher), FASTA, Intervals & Points, BMB, CLUSTAL V, CLUSTAL W, CONSENSUS, LCONSENSUS, WCONSENSUS, Smith-Waterman algorithm, DARWIN, Las Vegas algorithm, FNAT (Forced Nucleotide Alignment Tool), Framealign, Framesearch, DYNAMIC, FILTER, FSAP (Fristensky Sequence Analysis Package), GAP (Global Alignment Program), GENAL, GIBBS, GenQuest, ISSC (Sensitive Sequence Comparison), LALIGN (Local Sequence Alignment), LCP (Local Content Program), MACAW (Multiple Alignment Construction & Analysis Workbench), MAP (Multiple Alignment Program), MBLKP, MBLKN, PIMA (Pattern-Induced Multi-sequence Alignment), SAGA (Sequence Alignment by Genetic Algorithm) and WHAT-IF. Such alignment programs can also be used to screen genome databases to identify polynucleotide sequences having substantially identical sequences.

[0062] A number of genome databases are available for comparison. Several databases containing genomic information annotated with some functional information are maintained by different organizations, and are accessible via the internet, for example, at world wide web addresses (url's) "wwwtigr.org/tdb"; "genetics.wisc.edu"; "genome-www.stanford.edu/~ball"; "hiv-web.lanl.gov"; "ncbi.nlm.nih.gov"; "cbi.ac.uk"; "Pasteur.fr/other/biology"; and "genome-wi.mit.edu".

[0063] In particular, the BLAST and BLAST 2.0 algorithms using default parameters are particularly useful for identifying polynucleotide and polypeptides encompassed within the present invention (Altschul et al. (Nucleic Acids Res. 25:3389-3402, 1977; J. Mol. Biol. 215:403-410, 1990, each of which is incorporated herein by reference). Software for performing BLAST analyses is publicly available through the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov). This algorithm involves first identifying high scoring sequence pairs (HSPs) by identifying short words of length W in the query sequence, which either match or satisfy some positive-valued threshold score T when aligned with a word of the same length in a database sequence. T is referred to as the neighborhood word score threshold (Altschul et al., supra, 1977, 1990). These initial neighborhood word hits act as seeds for initiating searches to find longer HSPs containing them. The word hits are extended in both directions along each sequence for as far as the cumulative alignment score can be increased. Cumulative scores are calculated using, for nucleotide sequences, the parameters M (reward score for a pair of matching residues; always >0). For amino acid sequences, a scoring matrix is used to calculate the cumulative score. Extension of the word hits in each direction are halted when: the cumulative alignment score falls off by the quantity X from its maximum achieved value; the cumulative score goes to zero or below, due to the accumulation of one or more negative-scoring residue alignments; or the end of either sequence is reached. The BLAST algorithm parameters W, T, and X determine the

sensitivity and speed of the alignment. The BLASTN program (for nucleotide sequences) uses as defaults a wordlength (W) of 11, an expectation (E) of 10, M=5, N=4 and a comparison of both strands. For amino acid sequences, the BLASTP program uses as defaults a wordlength of 3, and expectations (E) of 10, and the BLOSUM62 scoring matrix (see Henikoff and Henikoff, <u>Proc. Natl. Acad. Sci., USA</u> 89:10915, 1989) alignments (B) of 50, expectation (E) of 10, M=5, N=4, and a comparison of both strands.

[0064] The BLAST algorithm also performs a statistical analysis of the similarity between two sequences (see, for example, Karlin and Altschul, Proc. Natl. Acad. Sci., USA 90:5873, 1993, which is incorporated herein by reference). One measure of similarity provided by BLAST algorithm is the smallest sum probability (P(N)), which provides an indication of the probability by which a match between two nucleotide or amino acid sequences would occur by chance. For example, a nucleic acid is considered similar to a references sequence if the smallest sum probability in a comparison of the test nucleic acid to the reference nucleic acid is less than about 0.2, more preferably less than about 0.01, and most preferably less than about 0.001. Significantly, upon identifying polynucleotides that are substantially similar to those of SEQ ID NOS:1-5379, the identified polynucleotides can be used as query sequences in a BLAST search to identify polynucleotides and polypeptides substantially similar thereto.

[0065] It should be noted that the nucleotide sequences set forth as SEQ ID NOS:1-2703 comprise coding sequences, whereas the nucleotide sequences set forth as SEQ ID NOS:2704-5379 comprise regulatory sequences. In addition, the coding sequences and regulatory sequences are related in that, for example, SEQ ID NO:1 is the coding sequence of a plant cold regulated gene having a 5' upstream (regulatory) sequence set forth as SEQ ID NO:2704 (see Table 2). Similarly, SEQ ID NO:2705 comprises a regulatory region of SEQ ID NO:2, SEQ ID NO:2706 comprises a regulatory region of SEQ ID NO:3, and so forth as shown in Table 2. As such, reference herein, for example, to a "polynucleotide comprising SEQ ID NO:1" can,

unless indicated otherwise, include at least SEQ ID NO:2704. In some cases, the entire coding region of a plant stress regulated gene or the 5' upstream sequence has not yet been determined (see, for example, SEQ ID NO:43 in Table 3, where "none" indicates that 5' upstream regulatory sequences have not yet been determined). However, the determination of a complete coding sequence where only a portion is known or of regulatory sequences where a portion of the coding sequence is known can be made using methods as disclosed herein or otherwise known in the art.

[0066] In one embodiment, protein and nucleic acid sequence homologies are evaluated using the Basic Local Alignment Search Tool ("BLAST"). In particular, five specific BLAST programs are used to perform the following task:

- BLASTP and BLAST3 compare an amino acid query sequence against a protein sequence database;
- (2) BLASTN compares a nucleotide query sequence against a nucleotide sequence database;
- (3) BLASTX compares the six-frame conceptual translation products of a query nucleotide sequence (both strands) against a protein sequence database;
- (4) TBLASTN compares a query protein sequence against a nucleotide sequence database translated in all six reading frames (both strands); and
- (5) TBLASTX compares the six-frame translations of a nucleotide query sequence against the six-frame translations of a nucleotide sequence database.
- [0067] The BLAST programs identify homologous sequences by identifying similar segments, which are referred to herein as "high-scoring segment pairs," between a query amino or nucleic acid sequence and a test sequence which is preferably obtained from a protein or nucleic acid sequence database. High-scoring segment pairs are preferably identified (i.e., aligned) by means of a scoring matrix, many of which are known in the art. Preferably, the scoring matrix used is the BLOSUM62 matrix (Gonnet et al., Science 256:1443-1445, 1992; Henikoff and Henikoff, Proteins 17:49-61, 1993, each of which is incorporated herein by

reference). Less preferably, the PAM or PAM250 matrices may also be used (Schwartz and Dayhoff, eds., "Matrices for Detecting Distance Relationships: Atlas of Protein Sequence and Structure" (Washington, National Biomedical Research Foundation 1978)). BLAST programs are accessible through the U.S. National Library of Medicine, for example, on the world wide web at address (url) "nebi.nlm.nih.gov".

[0068] The parameters used with the above algorithms may be adapted depending on the sequence length and degree of homology studied. In some embodiments, the parameters may be the default parameters used by the algorithms in the absence of instructions from the user.

[0069] The term "substantially similar" also is used in reference to a comparison of expression profiles of nucleotide sequences, wherein a determination that an expression profile characteristic of a stress response is substantially similar to the profile of nucleic acid molecules expressed in a plant cell being examined ("test plant") is indicative of exposure of the test plant cell to one or a combination of abiotic stress conditions. When used in reference to such a comparison of expression profiles, the term "substantially similar" means that that the individual nucleotide sequences in the test plant cell profile are altered in the same manner as the corresponding nucleotide sequences in the expression profile characteristic of the stress response.

[0070] By way of example, where exposure to saline results in an increased expression of nucleotide sequences A, B and C, and a decreased expression of nucleotide sequences D and E, as indicated by the expression profile characteristic of a saline stress response, a determination that corresponding nucleotide sequences A, B and C in the test plant cell are increased and that nucleotides sequences D and E are decreased is indicative of exposure of the test plant cell to a saline stress condition. It should be recognized that, where, for example, only nucleotide sequences A, B, D and E are examined in the test plant cell, an increase in A and B and a decrease in D and E

expression of the test plant cells is considered to be substantially similar to the expression profile characteristic of a saline stress condition and, therefore, is indicative of exposure of the plant cell to a saline stress condition. Similarly, where the levels of expression of the nucleotide sequences examined in a test plant are altered in the same manner, i.e., are increased or are decreased, as that observed in an expression profile characteristic of a particular stress response, the absolute levels of expression may vary, for example, two-fold, five-fold, ten-fold, or the like.

Nevertheless, the expression profile of the test plant cell is considered to be substantially similar to the expression profile characteristic of the particular stress response and, therefore, indicative of exposure of the plant cell to the stress condition.

[0071] As disclosed herein, clusters of stress-regulated genes (and their products), some of which also have been described as having cellular functions such as enzymatic activity or roles as transcription factors, are involved in the response of plant cells to various abiotic stresses (see Tables 29-31; see, also, Tables 1 and 32). As such, the polynucleotide sequences comprising the genes in a cluster likely share common stress-regulated regulatory elements, including, for example, cold-regulated regulatory elements (SEQ ID NOS:2704-3955), salinity-regulated regulatory elements (SEQ ID NOS:4910-5107, and osmotic pressure-regulated regulatory elements (SEQ ID NOS:5108-5263), as well as regulatory elements that are responsive to a combination of stress conditions, but not to any of the individual stress conditions, alone (SEQ ID NOS:3956-4909 and 5263-5379). The identification of such clusters of genes thus provides a means to identify the stress-regulated regulatory elements that control the level of expression of these genes.

[0072] As used herein, the term "plant stress-regulated gene" means a polynucleotide sequence of a plant, the transcription of which is altered in response to exposure to a stress condition, and the regulatory elements linked to such a polynucleotide sequence and involved in the stress response, which can be induction or repression. In general, plant stress gene regulatory elements are contained within a sequence including approximately two kilobases upstream (5') of the transcription or

translation start site and two kilobases downstream (3') of the transcription or translation termination site. In the absence of an abiotic stress condition, the stress-regulated gene can normally be unexpressed in the cells, can be expressed at a basal level, which is induced to a higher level in response to the stress condition, or can be expressed at a level that is reduced (decreased) in response to the stress condition. The coding region of a plant stress-regulated gene encodes a stress-regulated polypeptide, and also can be the basis for expression of a functional RNA molecule such as an antisense molecule or ribozyme. A stress-regulated polypeptide can have an adaptive effect on a plant, thereby allowing the plant to better tolerate stress conditions; or can have a maladaptive effect, thereby decreasing the ability of the plant to tolerate the stress conditions.

The present invention provides an isolated plant stress-regulated regulatory 100731 element, which regulates expression of an operatively linked nucleotide sequence in a plant in response a stress condition. As disclosed herein, a plant stress-regulated regulatory element can be isolated from a polynucleotide sequence of a plant stressregulated gene comprising a nucleotide sequence as set forth in SEQ ID NOS:1-2703, for example any of SEQ ID NOS:2704-5379 (see Table 2). It is recognized that certain of the polynucleotides set forth as SEQ ID NOS:1-5379 previously have been described as being involved in a stress-regulated response in plants, including SEQ ID NOS:156, 229, 233, 558, 573, 606, 625, 635, 787, 813, 1263, 1386, 1391, 1405, 1445, 1484, 1589, 1609, 1634, 1726, 1866, 1918, and 1928 and, therefore, are not encompassed, in whole or in part, within the compositions of the invention, and are encompassed within only certain particular methods of the invention, for example, methods of making a transgenic plant that is resistant to two or more stress conditions, since, even where such a gene was known to be expressed in response to a single stress condition such as cold or saline (e.g., SEQ ID NO:1263), it was not known prior to the present disclosure that any of these genes was responsive to a combination of stress conditions (for example, a combination of cold and osmotic stress for SEQ ID NOS:1726, 1866, 1918, and 1928; or a combination of cold, osmotic and saline stress for SEQ ID NOS:1263,1386, 1391, 1405, 1445, 1484, 1589, 1609, and 1634).

[0074] Methods for identifying and isolating the stress-regulated regulatory element from the disclosed polynucleotides, or genomic DNA clones corresponding thereto, are well known in the art. For example, methods of making deletion constructs or linker-scanner constructs can be used to identify nucleotide sequences that are responsive to a stress condition. Generally, such constructs include a reporter gene operatively linked to the sequence to be examined for regulatory activity. By performing such assays, a plant stress-regulated regulatory element can be defined within a sequence of about 500 nucleotides or fewer, generally at least about 200 nucleotides or fewer, particularly about 50 to 100 nucleotides, and more particularly at least about 20 nucleotides or fewer. Preferably the minimal (core) sequence required for regulating a stress response of a plant is identified.

[0075] The nucleotide sequences of the genes of a cluster also can be examined using a homology search engine such as described herein to identify sequences of conserved identity, particularly in the nucleotide sequence upstream of the transcription start site. Since all of the genes in a cluster as disclosed are induced in response to a particular stress condition or a particular combination of stress conditions, some or all of the nucleotide sequences can share conserved stress-regulated regulatory elements. By performing such a homology search, putative stress-regulated regulatory elements can be identified. The ability of such identified sequences to function as a plant stress-regulated regulatory element can be confirmed, for example, by operatively linking the sequence to a reporter gene and assaying the construct for responsiveness to a stress condition.

[0076] As used herein, the term "regulatory element" means a nucleotide sequence that, when operatively linked to a coding region of a gene, effects transcription of the coding region such that a ribonucleic acid (RNA) molecule is transcribed from the coding region. A regulatory element generally can increase or decrease the amount of transcription of a nucleotide sequence, for example, a coding sequence, operatively linked to the element with respect to the level at which the nucleotide sequence would be transcribed absent the regulatory element. Regulatory elements are well known in

the art and include promoters, enhancers, silencers, inactivated silencer intron sequences, 3'-untranslated or 5'-untranslated sequences of transcribed sequence, for example, a poly-A signal sequence, or other protein or RNA stabilizing elements, or other gene expression control elements known to regulate gene expression or the amount of expression of a gene product. A regulatory element can be isolated from a naturally occurring genomic DNA sequence or can be synthetic, for example, a synthetic promoter.

[0077] Regulatory elements can be constitutively expressed regulatory element, which maintain gene expression at a relative level of activity (basal level), or can be regulated regulatory elements. Constitutively expressed regulatory elements can be expressed in any cell type, or can be tissue specific, which are expressed only in particular cell types, phase specific, which are expressed only during particular developmental or growth stages of a plant cell, or the like. A regulatory element such as a tissue specific or phase specific regulatory element or an inducible regulatory element useful in constructing a recombinant polynucleotide or in a practicing a method of the invention can be a regulatory element that generally, in nature, is found in a plant genome. However, the regulatory element also can be from an organism other than a plant, including, for example, from a plant virus, an animal virus, or a cell from an animal or other multicellular organism.

[0078] A regulatory element useful for practicing method of the present is a promoter element. Useful promoters include, but are not limited to, constitutive, inducible, temporally regulated, developmentally regulated, spatially-regulated, chemically regulated, stress-responsive, tissue-specific, viral and synthetic promoters. Promoter sequences are known to be strong or weak. A strong promoter provides for a high level of gene expression, whereas a weak promoter provides for a very low level of gene expression. An inducible promoter is a promoter that provides for the turning on and off of gene expression in response to an exogenously added agent, or to an environmental or developmental stimulus. A bacterial promoter such as the Ptace promoter can be induced to varying levels of gene expression depending on the level

of isothiopropylgalactoside added to the transformed bacterial cells. An isolated promoter sequence that is a strong promoter for heterologous nucleic acid is advantageous because it provides for a sufficient level of gene expression to allow for easy detection and selection of transformed cells and provides for a high level of gene expression when desired.

[0079] Within a plant promoter region there are several domains that are necessary for full function of the promoter. The first of these domains lies immediately upstream of the structural gene and forms the "core promoter region" containing consensus sequences, normally 70 base pairs immediately upstream of the gene. The core promoter region contains the characteristic CAAT and TATA boxes plus surrounding sequences, and represents a transcription initiation sequence that defines the transcription start point for the structural gene.

[0080] The presence of the core promoter region defines a sequence as being a promoter: if the region is absent, the promoter is non-functional. The core promoter region, however, is insufficient to provide full promoter activity. A series of regulatory sequences upstream of the core constitute the remainder of the promoter. These regulatory sequences determine expression level, the spatial and temporal pattern of expression and, for an important subset of promoters, expression under inductive conditions (regulation by external factors such as light, temperature, chemicals, hormones).

[0081] To define a minimal promoter region, a DNA segment representing the promoter region is removed from the 5' region of the gene of interest and operably linked to the coding sequence of a marker (reporter) gene by recombinant DNA techniques well known to the art. The reporter gene is operably linked downstream of the promoter, so that transcripts initiating at the promoter proceed through the reporter gene. Reporter genes generally encode proteins which are easily measured, including, but not limited to, chloramphenicol acetyl transferase (CAT), beta-glucuronidase (GUS), green fluorescent protein (GFP), β-galactosidase (β-GAL), and luciferase.

[0082] The construct containing the reporter gene under the control of the promoter is then introduced into an appropriate cell type by transfection techniques well known to the art. To assay for the reporter protein, cell lysates are prepared and appropriate assays, which are well known in the art, for the reporter protein are performed. For example, if CAT were the reporter gene of choice, the lysates from cells transfected with constructs containing CAT under the control of a promoter under study are mixed with isotopically labeled chloramphenicol and acetyl-coenzyme A (acetyl-CoA). The CAT enzyme transfers the acetyl group from acetyl-CoA to the 2-position or 3-position of chloramphenicol. The reaction is monitored by thin layer chromatography, which separates acetylated chloramphenicol from unreacted material. The reaction products are then visualized by autoradiography.

[0083] The level of enzyme activity corresponds to the amount of enzyme that was made, which in turn reveals the level of expression from the promoter of interest. This level of expression can be compared to other promoters to determine the relative strength of the promoter under study. In order to be sure that the level of expression is determined by the promoter, rather than by the stability of the mRNA, the level of the reporter mRNA can be measured directly, for example, by northern blot analysis.

[0084] Once activity is detected, mutational and/or deletional analyses may be employed to determine the minimal region and/or sequences required to initiate transcription. Thus, sequences can be deleted at the 5' end of the promoter region and/or at the 3' end of the promoter region, and nucleotide substitutions introduced. These constructs are then introduced to cells and their activity determined.

[0085] The choice of promoter will vary depending on the temporal and spatial requirements for expression, and also depending on the target species. In some cases, expression in multiple tissues is desirable. While in others, tissue-specific, e.g., leaf-specific, seed-specific, petal-specific, anther-specific, or pith-specific, expression is desirable. Although many promoters from dicotyledons have been shown to be operational in monocotyledons and vice versa, ideally dicotyledonous promoters are

selected for expression in dicotyledons, and monocotyledonous promoters for expression in monocotyledons. There is, however, no restriction to the origin or source of a selected promoter. It is sufficient that the promoters are operational in driving the expression of a desired nucleotide sequence in the particular cell.

[0086] A range of naturally-occurring promoters are known to be operative in plants and have been used to drive the expression of heterologous (both foreign and endogenous) genes and nucleotide sequences in plants: for example, the constitutive 35S cauliflower mosaic virus (CaMV) promoter, the ripening-enhanced tomato polygalacturonase promoter (Bird et al., 1988), the E8 promoter (Diekman and Fischer, 1988) and the fruit specific 2A1 promoter (Pear et al., 1989). Many other promoters, e.g., U2 and U5 snRNA promoters from maize, the promoter from alcohol dehydrogenase, the Z4 promoter from a gene encoding the Z4 22 kD zein protein, the Z10 promoter from a gene encoding a 10 kD zein protein, a Z27 promoter from a gene encoding a 27 kD zein protein, the A20 promoter from the gene encoding a 19 kD zein protein, inducible promoters, such as the light inducible promoter derived from the pea rbcS gene and the actin promoter from rice, e.g., the actin 2 promoter (WO 00/70067); seed specific promoters, such as the phaseolin promoter from beans, may also be used. The nucleotide sequences of the stress-regulated genes of this invention can also be expressed under the regulation of promoters that are chemically regulated. This enables the nucleic acid sequence or encoded polypeptide to be synthesized only when the crop plants are treated with the inducing chemicals. Chemical induction of gene expression is detailed in EP 0 332 104 and U.S. Pat. 5.614,395.

[0087] In some instances it may be desirable to link a constitutive promoter to a polynucleotide comprising a stress regulated gene of the invention. Examples of some constitutive promoters include the rice actin 1 (Wang et al., 1992; U.S. Pat. No. 5,641,876), CaMV 35S (Odeil et al., 1985), CaMV 19S (Lawton et al., 1987), nos. Adh. sucrose synthase: and the ubiquitin promoters.

[0088] In other situations it may be desirable to limit expression of stress-related sequences to specific tissues or stages of development. As used herein, the term "tissue specific or phase specific regulatory element" means a nucleotide sequence that effects transcription in only one or a few cell types, or only during one or a few stages of the life cycle of a plant, for example, only for a period of time during a particular stage of growth, development or differentiation. The terms "tissue specific" and "phase specific" are used together herein in referring to a regulatory element because a single regulatory element can have characteristics of both types of regulatory elements. For example, a regulatory element active only during a particular stage of plant development also can be expressed only in one or a few types of cells in the plant during the particular stage of development. As such, any attempt to classify such regulatory elements as tissue specific or as phase specific can be difficult. Accordingly, unless indicated otherwise, all regulatory elements having the characteristic of a tissue specific regulatory element, or a phase specific regulatory element, or both are considered together for purposes of the present invention.

[0089] Examples of tissue specific promoters which have been described include the lectin (Vodkin, 1983; Lindstrom et al., 1990) corn alcohol dehydrogenase 1 (Vogel et al., 1989; Dennis et al., 1984), corn light harvesting complex (Simpson, 1986; Bansal et al., 1992), corn heat shock protein (Odell et al., 1985), pea small subunit RuBP carboxylase (Poulsen et al., 1986), Ti plasmid mannopine synthase and Ti plasmid nopaline synthase (Langridge et al., 1989), petunia chalcone isomerase (vanTunen et al., 1988), bean glycine rich protein 1 (Keller et al., 1989), truncated CaMV 35s (Odell et al., 1985), potato patatin (Wenzler et al., 1989), root cell (Yamamoto et al., 1990), maize zein (Reina et al., 1990; Kriz et al., 1987; Wandelt et al., 1989; Langridge et al., 1983; Reina et al., 1990), globulin-1 (Belanger et al., 1991), α-tubulin, cab (Sullivan et al., 1989), PEPCase (Hudspeth & Grula, 1989), R gene complex-associated promoters (Chandler et al., 1989), histone, and chalcone synthase promoters (Franken et al., 1991). Tissue specific enhancers are described by Fromm et al. (1989).

[0090] Several other tissue-specific regulated genes and/or promoters have been reported in plants, including genes encoding seed storage proteins such as napin, cruciferin, beta-conglycinin, and phaseolin, zein or oil body proteins such as oleosin, genes involved in fatty acid biosynthesis, including acyl carrier protein, stearoyl-ACP desaturase, fatty acid desaturases (fad 2-1), and other genes expressed during embryonic development such as Bce4 (see, for example, EP 255378 and Kridl et al., 1991). Particularly useful for seed-specific expression is the pea vicilin promoter (Czako et al., 1992). (See also U.S. Pat. No. 5,625,136, which is incorporated herein by reference.) Other useful promoters for expression in mature leaves are those that are switched on at the onset of senescence, such as the SAG promoter from Arabidopsis (Gan et al., 1995).

[0091] A class of fruit-specific promoters expressed at or during antithesis through fruit development, at least until the beginning of ripening, is discussed in U.S. Pat. No. 4,943,674. cDNA clones that are preferentially expressed in cotton fiber have been isolated (John et al., 1992). cDNA clones from tomato displaying differential expression during fruit development have been isolated and characterized (Mansson et al., 1985, Slater et al., 1985). The promoter for polygalacturonase gene is active in fruit ripening. The polygalacturonase gene is described in U.S. Pat. Nos. 4,535,060, 4,769,061, 4,801,590, and 5,107,065, each of which is incorporated herein by reference.

[0092] Other examples of tissue-specific promoters include those that direct expression in leaf cells following damage to the leaf (for example, from chewing insects), in tubers (for example, patatin gene promoter), and in fiber cells (an example of a developmentally-regulated fiber cell protein is E6 (John et al., 1992). The E6 gene is most active in fiber, although low levels of transcripts are found in leaf, oxylle and flower.

[0093] Additional tissue specific or phase specific regulatory elements include, for example, the AGL8/FRUITFULL regulatory element, which is activated upon floral

induction (Hempel et al., Development 124:3845-3853, 1997, which is incorporated herein by reference); root specific regulatory elements such as the regulatory elements from the RCP1 gene and the LRP1 gene (Tsugeki and Fedoroff, Proc. Natl. Acad., USA 96:12941-12946, 1999; Smith and Fedoroff, Plant Cell 7:735-745, 1995, each of which is incorporated herein by reference); flower specific regulatory elements such as the regulatory elements from the LEAFY gene and the APETELA1 gene (Blazquez et al., Development 124:3835-3844, 1997, which is incorporated herein by reference; Hempel et al., supra, 1997); seed specific regulatory elements such as the regulatory element from the oleosin gene (Plant et al., Plant Mol. Biol. 25:193-205, 1994, which is incorporated herein by reference), and dehiscence zone specific regulatory element. Additional tissue specific or phase specific regulatory elements include the Zn13 promoter, which is a pollen specific promoter (Hamilton et al., Plant Mol. Biol. 18:211-218, 1992, which is incorporated herein by reference); the UNUSUAL FLORAL ORGANS (UFO) promoter, which is active in apical shoot meristem; the promoter active in shoot meristems (Atanassova et al., Plant J. 2:291, 1992, which is incorporated herein by reference), the cdc2a promoter and cyc07 promoter (see, for example. Ito et al., Plant Mol. Biol. 24:863, 1994; Martinez et al., Proc. Natl. Acad. Sci., USA 89:7360, 1992; Medford et al., Plant Cell 3:359, 1991; Terada et al., Plant J. 3:241, 1993; Wissenbach et al., Plant J. 4:411, 1993, each of which is incorporated herein by reference); the promoter of the APETELA3 gene, which is active in floral meristems (Jack et al., Cell 76:703, 1994, which is incorporated herein by reference; Hempel et al., supra, 1997); a promoter of an agamous-like (AGL) family member, for example, AGL8, which is active in shoot meristem upon the transition to flowering (Hempel et al., supra, 1997); floral abscission zone promoters; L1-specific promoters; and the like.

[0094] The tissue-specificity of some "tissue-specific" promoters may not be absolute and may be tested by one skilled in the art using the diphtheria toxin sequence. One can also achieve tissue-specific expression with "leaky" expression by a combination of different tissue-specific promoters (Beals et al., 1997). Other tissue-specific promoters can be isolated by one skilled in the art (see U.S. 5,589,379).

Several inducible promoters ("gene switches") have been reported, many of which are described in the review by Gatz (1996) and Gatz (1997). These include tetracycline repressor system, *Lac* repressor system, copper inducible systems, salicylate inducible systems (such as the PR1a system), glucocorticoid (Aoyama et al., 1997) and ecdysone inducible systems. Also included are the benzene sulphonamide (U.S. Pat. No. 5,364,780) and alcohol (WO 97/06269 and WO 97/06268) inducible systems and glutathione S-transferase promoters.

[0095] In some instances it might be desirable to inhibit expression of a native DNA sequence within a plant's tissues to achieve a desired phenotype. In this case, such inhibition might be accomplished with transformation of the plant to comprise a constitutive, tissue-independent promoter operably linked to an antisense nucleotide sequence, such that constitutive expression of the antisense sequence produces an RNA transcript that interferes with translation of the mRNA of the native DNA sequence.

[0096] Inducible regulatory elements also are useful for purposes of the present invention. As used herein, the term "inducible regulatory element" means a regulatory element that, when exposed to an inducing agent, effects an increased level of transcription of a nucleotide sequence to which it is operatively linked as compared to the level of transcription, if any, in the absence of an inducing agent. Inducible regulatory elements can be those that have no basal or constitutive activity and only effect transcription upon exposure to an inducing agent, or those that effect a basal or constitutive level of transcription, which is increased upon exposure to an inducing agent. Inducible regulatory elements that effect a basal or constitutive level of expression generally are useful in a method or composition of the invention where the induced level of transcription is substantially greater than the basal or constitutive level of expression, for example, at least about two-fold greater, or at least about five-fold greater. Particularly useful inducible regulatory elements do not have a basal or constitutive activity, or increase the level of transcription at least about ten-fold

greater than a basal or constitutive level of transcription associated with the regulatory element.

[0097] Inducible promoters that have been described include the ABA- and turgor-inducible promoters, the promoter of the auxin-binding protein gene (Schwob et al., 1993), the UDP glucose flavonoid glycosyl-transferase gene promoter (Ralston et al., 1988), the MPI proteinase inhibitor promoter (Cordero et al., 1994), and the glyceraldehyde-3-phosphate dehydrogenase gene promoter (Kohler et al., 1995; Quigley et al., 1989; Martinez et al., 1989).

[0098] The term "inducing agent" is used to refer to a chemical, biological or physical agent or environmental condition that effects transcription from an inducible regulatory element. In response to exposure to an inducing agent, transcription from the inducible regulatory element generally is initiated *de novo* or is increased above a basal or constitutive level of expression. Such induction can be identified using the methods disclosed herein, including detecting an increased level of RNA transcribed from a nucleotide sequence operatively linked to the regulatory element, increased expression of a polypeptide encoded by the nucleotide sequence, or a phenotype conferred by expression of the encoded polypeptide.

[0099] An inducing agent useful in a method of the invention is selected based on the particular inducible regulatory element. For example, the inducible regulatory element can be a metallothionein regulatory element, a copper inducible regulatory element or a tetracycline inducible regulatory element, the transcription from which can be effected in response to metal ions, copper or tetracycline, respectively (Furst et al., Cell 55:705-717, 1988; Mett et al., Proc. Natl. Acad. Sci., USA 90:4567-4571, 1993; Gatz et al., Plant J. 2:397-404, 1992; Roder et al., Mol. Gen. Genet. 243:32-38, 1994, each of which is incorporated herein by reference). The inducible regulatory element also can be an ecdysone regulatory element or a glucocorticoid regulatory element, the transcription from which can be effected in response to ecdysone or other steroid (Christopherson et al., Proc. Natl. Acad. Sci., USA 89:6314-6318, 1992;

Schena et al., <u>Proc. Natl. Acad. Sci., USA</u> 88:10421-10425, 1991, each of which is incorporated herein by reference). In addition, the regulatory element can be a cold responsive regulatory element or a heat shock regulatory element, the transcription of which can be effected in response to exposure to cold or heat, respectively (Takahashi et al., <u>Plant Physiol.</u> 99:383-390, 1992, which is incorporated herein by reference). Additional regulatory elements useful in the methods or compositions of the invention include, for example, the spinach nitrite reductase gene regulatory element (Back et al., <u>Plant Mol. Biol.</u> 17:9, 1991, which is incorporated herein by reference); a light inducible regulatory element (Feinbaum et al., <u>Mol. Gen. Genet.</u> 226:449, 1991; Lam and Chua, <u>Science</u> 248:471, 1990, each of which is incorporated herein by reference), a plant hormone inducible regulatory element (Yamaguchi-Shinozaki et al., <u>Plant Mol. Biol.</u> 15:905, 1990; Kares et al., <u>Plant Mol. Biol.</u> 15:225, 1990, each of which is incorporated herein by reference), and the like.

[0100] An inducible regulatory element also can be a plant stress-regulated regulatory element of the invention. In addition to the known stress conditions that specifically induce or repress expression from such elements, the present invention provides methods of identifying agents that mimic a stress condition. Accordingly, such stress mimics are considered inducing or repressing agents with respect to a plant stress-regulated regulatory element. In addition, a recombinant polypeptide comprising a zinc finger domain, which is specific for the regulatory element, and an effector domain, particularly an activator, can be useful as an inducing agent for a plant stress-regulated regulatory element. Furthermore, such a recombinant polypeptide provides the advantage that the effector domain can be a repressor domain, thereby providing a repressing agent, which decreases expression from the regulatory element. In addition, use of such a method of modulating expression of an endogenous plant stress-regulated gene provides the advantage that the polynucleotide encoding the recombinant polypeptide can be introduced into cells of the plant, thus providing a transgenic plant that can be regulated coordinately with the endogenous plant stress-regulated gene upon exposure to a stress condition. A polynucleotide encoding such a recombinant polypeptide can be operatively linked to and expressed

from a constitutively active, inducible or tissue specific or phase specific regulatory element.

[0101] In one embodiment, the promoter may be a gamma zein promoter, an oleosin ole16 promoter, a globulin I promoter, an actin I promoter, an actin cl promoter, a sucrose synthetase promoter, an INOPS promoter, an EXM5 promoter, a globulin2 promoter, a b-32, ADPG-pyrophosphorylase promoter, an LtpI promoter, an Ltp2 promoter, an oleosin ole17 promoter, an oleosin ole18 promoter, an actin 2 promoter, a pollen-specific protein promoter, a pollen-specific pectate lyase promoter, an anther-specific protein promoter (Huffman), an anther-specific gene RTS2 promoter, a pollen-specific gene promoter, a tapeturn-specific gene promoter, taneturn- specific gene RAB24 promoter, a anthranilate synthase alpha subunit promoter, an alpha zein promoter, an anthranilate synthase beta subunit promoter, a dihydrodipicolinate synthase promoter, a Thi l promoter, an alcohol dehydrogenase promoter, a cab binding protein promoter, an H3C4 promoter, a RUBISCO SS starch branching enzyme promoter, an ACCase promoter, an actin3 promoter, an actin7 promoter, a regulatory protein GF14-12 promoter, a ribosomal protein L9 promoter, a cellulose biosynthetic enzyme promoter, an S-adenosyl-L-homocysteine hydrolase promoter, a superoxide dismutase promoter, a C-kinase receptor promoter, a phosphoglycerate mutase promoter, a root-specific RCc3 mRNA promoter, a glucose-6 phosphate isomerase promoter, a pyrophosphate-fructose 6-phosphate-lphosphotransferase promoter, an ubiquitin promoter, a beta-ketoacyl-ACP synthase promoter, a 33 kDa photosystem 11 promoter, an oxygen evolving protein promoter, a 69 kDa vacuolar ATPase subunit promoter, a metallothionein-like protein promoter, a glyceraldehyde-3-phosphate dehydrogenase promoter, an ABA- and ripeninginducible-like protein promoter, a phenylalanine ammonia lyase promoter, an adenosine triphosphatase S-adenosyl-L-homocysteine hydrolase promoter, an atubulin promoter, a cab promoter, a PEPCase promoter, an R gene promoter, a lectin promoter, a light harvesting complex promoter, a heat shock protein promoter, a chalcone synthase promoter, a zein promoter, a globulin-1 promoter, an ABA promoter, an auxin-binding protein promoter, a UDP glucose flavonoid glycosyltransferase gene promoter, an NTI promoter, an actin promoter, an opaque 2 promoter, a b70 promoter, an oleosin promoter, a CaMV 35S promoter, a CaMV 19S promoter, a histone promoter, a turgor-inducible promoter, a pea small subunit RuBP carboxylase promoter, a Ti plasmid mannopine synthase promoter, Ti plasmid nopaline synthase promoter, a petunia chalcone isomerase promoter, a bean glycine rich protein I promoter, a CaMV 35S transcript promoter, a potato patatin promoter, or a S-E9 small subunit RuBP carboxylase promoter.

[0102] In addition to promoters, a variety of 5' and 3' transcriptional regulatory sequences are also available for use in the present invention. Transcriptional terminators are responsible for the termination of transcription and correct mRNA polyadenylation. The 3'-untranslated regulatory DNA sequence preferably includes from about 50 to about 1,000, more preferably about 100 to about 1,000, nucleotide base pairs and contains plant transcriptional and translational termination sequences. Appropriate transcriptional terminators and those which are known to function in plants include the CaMV 35S terminator, the tml terminator, the nopaline synthase terminator, the pea rbcS E9 terminator, the terminator for the T7 transcript from the octopine synthase gene of Agrobacterium tumefaciens, and the 3' end of the protease inhibitor I or II genes from potato or tomato, although other 3' elements known to those of skill in the art can also be employed. Alternatively, one also could use a gamma coixin, oleosin 3 or other terminator from the genus Coix. Preferred 3' elements include those from the nopaline synthase gene of Agrobacterium tumefaciens (Bevan et al., 1983), the terminator for the T7 transcript from the octopine synthase gene of Agrobacterium tumefaciens, and the 3' end of the protease inhibitor I or II genes from potato or tomato.

[0103] As the DNA sequence between the transcription initiation site and the start of the coding sequence, i.e., the untranslated leader sequence, can influence gene expression, one may also wish to employ a particular leader sequence. Preferred leader sequences are contemplated to include those that include sequences predicted to direct optimum expression of the attached sequence, i.e., to include a preferred

consensus leader sequence that may increase or maintain mRNA stability and prevent inappropriate initiation of translation. The choice of such sequences will be known to those of skill in the art in light of the present disclosure. Sequences that are derived from genes that are highly expressed in plants will be most preferred.

[0104] Other sequences that have been found to enhance gene expression in transgenic plants include intron sequences (e.g., from Adhl, bronzel, actinl, actin 2 (WO 00/760067), or the sucrose synthase intron) and viral leader sequences (e.g., from TMV, MCMV and AMV). For example, a number of non-translated leader sequences derived from viruses are known to enhance expression. Specifically, leader sequences from tobacco mosaic virus (TMV), maize chlorotic mottle virus (MCMV), and alfalfa mosaic virus (AMV) have been shown to be effective in enhancing expression (e.g., Gallie et al., 1987; Skuzeski et al., 1990). Other leaders known in the art include but are not limited to picornavirus leaders, for example, EMCV leader (encephalomyocarditis virus 5' non-coding region; Elroy-Stein et al., 1989); potyvirus leaders, for example, TEV leader (tobacco etch virus); MDMV leader (maize dwarf mosaic virus); human immunoglobulin heavy chain binding protein (BiP) leader, (Macejak et al., 1991); untranslated leader from the coat protein mRNA of AMV (AMV RNA 4; Jobling et al., 1987), TMV (Gallie et al., 1989), and MCMV (Lommel et al., 1991; see also, della Cioppa et al., 1987).

[0105] Regulatory elements such as Adh intron 1 (Callis et al., 1987), sucrose synthase intron (Vasil et al., 1989) or TMV omega element (Gallie, et al., 1989), may further be included where desired. Examples of enhancers include elements from the CaMV 35S promoter, octopine synthase genes (Ellis et al., 1987), the rice actin I gene, the maize alcohol dehydrogenase gene (Callis et al., 1987), the maize shrunken I gene (Vasil et al., 1989), TMV Omega element (Gallie et al., 1989) and promoters from non-plant eukaryotes (e.g., yeast; Ma et al., 1988).

[0106] Vectors for use in accordance with the present invention may be constructed to include the ocs enhancer element, which was first identified as a 16 bp palindromic enhancer from the octopine synthase (ocs) gene of ultilane (Ellis et al., 1987), and is present in at least 10 other promoters (Bouchez et al., 1989). The use of an enhancer element, such as the ocs element and particularly multiple copies of the element, will act to increase the level of transcription from adjacent promoters when applied in the context of monocot transformation.

[0107] The methods of the invention provide genetically modified plant cells. which can contain, for example, a coding region, or peptide portion thereof, of a plant stress-regulated gene operatively linked to a heterologous inducible regulatory element; or a plant stress-regulated regulatory element operatively linked to a heterologous nucleotide sequence encoding a polypeptide of interest. In such a plant, the expression from the inducible regulatory element can be effected by exposing the plant cells to an inducing agent in any of numerous ways depending, for example, on the inducible regulatory element and the inducing agent. For example, where the inducible regulatory element is a cold responsive regulatory element present in the cells of a transgenic plant, the plant can be exposed to cold conditions, which can be produced artificially, for example, by placing the plant in a thermostatically controlled room, or naturally, for example, by planting the plant in an environment characterized, at least in part, by attaining temperatures sufficient to induce transcription from the promoter but not so cold as to kill the plants. By examining the phenotype of such transgenic plants, those plants that ectopically express a gene product that confers increased resistance of the plant to cold can be identified. Similarly, a transgenic plant containing a metallothionein promoter can be exposed to metal ions such as cadmium or copper by watering the plants with a solution containing the inducing metal ions, or can be planted in soil that is contaminated with a level of such metal ions that is toxic to most plants. The phenotype of surviving plants can be observed, those expressing desirable traits can be selected.

[0108] As used herein, the term "phenotype" refers to a physically detectable characteristic. A phenotype can be identified visually by inspecting the physical appearance of a plant following exposure, for example, to increased osmotic

conditions; can be identified using an assay to detecting a product produced due to expression of reporter gene, for example, an RNA molecule, a polypeptide such as an enzyme, or other detectable signal such as disclosed herein; or by using any appropriate tool useful for identifying a phenotype of a plant, for example, a microscope, a fluorescence activated cell sorter, or the like.

[0109] A transgenic plant containing an inducible regulatory element such as a steroid inducible regulatory element can be exposed to a steroid by watering the plants with a solution containing the steroid. The use of an inducible regulatory element that is induced upon exposure to a chemical or biological inducing agent that can be placed in solution or suspension in an aqueous medium can be particularly useful because the inducing agent can be applied conveniently to a relatively large crop of transgenic plants containing the inducible regulatory element, for example, through a watering system or by spraying the inducing agent over the field. As such, inducible regulatory elements that are responsive to an environmental inducing agent, for example, cold; heat; metal ions or other potentially toxic agents such as a pesticides, which can contaminate a soil; or the like; or inducible regulatory elements that are regulated by inducing agents that conveniently can be applied to plants, can be particularly useful in a method or composition of the invention, and allow the identification and selection of plants that express desirable traits and survive and grow in environments that otherwise would not support growth of the plants.

[0110] As disclosed herein, the present invention provides plant stress-regulated regulatory elements, which are identified based on the expression of clusters of plant genes in response to stress. As used herein, the term "stress-regulated regulatory element of a plant" or "plant stress-regulated regulatory element" means a nucleotide sequence of a plant genome that can respond to a stress such that expression of a gene product encoded by a gene comprising the regulatory element (a stress-inducible gene) is increased above or decreased below the level of expression of the gene product in the absence of the stress condition. The regulatory element can be any gene regulatory element, including, for example, a promoter, an enhancer, a silencer,

or the like. In one embodiment, the plant stress-regulated regulatory element is a plant stress-regulated promoter.

[0111] For purposes of modulating the responsiveness of a plant to a stress condition, it can be useful to introduce a modified plant stress-regulated regulatory element into a plant. Such a modified regulatory element can have any desirable characteristic, for example, it can be inducible to a greater level than the corresponding wild-type promoter, or it can be inactivated such that, upon exposure to a stress, there is little or no induction of expression of a nucleotide sequence operatively linked to the mutant element. A plant stress-regulated regulatory element can be modified by incorporating random mutations using, for example, *in vitro* recombination or DNA shuffling (Stemmer et al., Nature 370: 389-391, 1994; U.S. Pat. No. 5,605,793, each of which is incorporated herein by reference). Using such a method, millions of mutant copies of the polynucleotide, for example, stress-regulated regulatory element, can be produced based on the original nucleotide sequence, and variants with improved properties, such as increased inducibility can be recovered.

[0112] A mutation method such as DNA shuffling encompasses forming a mutagenized double-stranded polynucleotide from a template double-stranded polynucleotide, wherein the template double-stranded polynucleotide has been cleaved into double stranded random fragments of a desired size, and comprises the steps of adding to the resultant population of double-stranded random fragments one or more single or double stranded oligonucleotides, wherein the oligonucleotides comprise an area of identity and an area of heterology to the double stranded template polynucleotide; denaturing the resultant mixture of double stranded random fragments and oligonucleotides into single stranded fragments; incubating the resultant population of single stranded fragments with a polymerase under conditions that result in the annealing of the single stranded fragments at the areas of identity to form pairs of annealed fragments, the areas of identity being sufficient for one member of a pair to prime replication of the other, thereby forming a mutagenized double-stranded polynucleotide; and repeating the second and third steps for at least two further

cycles, wherein the resultant mixture in the second step of a further cycle includes the mutagenized double-stranded polynucleotide from the third step of the previous cycle, and the further cycle forms a further mutagenized double-stranded polynucleotide. Preferably, the concentration of a single species of double stranded random fragment in the population of double stranded random fragments is less than 1% by weight of the total DNA. In addition, the template double stranded polynucleotide can comprise at least about 100 species of polynucleotides. The size of the double stranded random fragments can be from about 5 base pairs to 5 kilobase pairs. In a further embodiment, the fourth step of the method comprises repeating the second and the third steps for at least 10 cycles.

[0113] A plant stress-regulated regulatory element of the invention is useful for expressing a nucleotide sequence operatively linked to the element in a cell, particularly a plant cell. As used herein, the term "expression" refers to the transcription and/or translation of an endogenous gene or a transgene in plants. In the case of an antisense molecule, for example, the term "expression" refers to the transcription of the polynucleotide encoding the antisense molecule.

[0114] As used herein, the term "operatively linked," when used in reference to a plant stress-regulated regulatory element, means that the regulatory element is positioned with respect to a second nucleotide sequence such that the regulatory element effects transcription or transcription and translation of the nucleotide sequence in substantially the same manner, but not necessarily to the same extent, as it does when the regulatory element is present in its natural position in a genome. Transcriptional promoters, for example, generally act in a position and orientation dependent manner and usually are positioned at or within about five nucleotides to about fifty nucleotides 5' (upstream) of the start site of transcription of a gene in nature. In comparison, enhancers and silencers can act in a relatively position or orientation independent manner and, therefore, can be positioned several hundred or thousand nucleotides upstream or downstream from a transcription start site, or in an

intron within the coding region of a gene, yet still be operatively linked to a coding region so as to effect transcription.

The second nucleotide sequence, i.e., the sequence operatively linked to the plant stress-regulated regulatory element, can be any nucleotide sequence, including, for example, a coding region of a gene or cDNA; a sequence encoding an antisense molecule, an RNAi molecule, ribozyme, triplexing agent (see, for example, Frank-Kamenetskii and Mirkin, Ann. Rev. Biochem. 64:65-95, 1995), or the like; or a sequence that, when transcribed, can be detected in the cell using, for example, by hybridization or amplification, or when translated produces a detectable signal. The term "coding region" is used broadly herein to include a nucleotide sequence of a genomic DNA or a cDNA molecule comprising all or part of a coding region of the coding strand. A coding region can be transcribed from an operatively linked regulatory element, and can be translated into a full length polypeptide or a peptide portion of a polypeptide. It should be recognized that, in a nucleotide sequence comprising a coding region, not all of the nucleotides in the sequence need necessarily encode the polypeptide and, particularly, that a gene transcript can contain one or more introns, which do not encode an amino acid sequence of a polypeptide but, nevertheless, are part of the coding region, particularly the coding strand, of the gene.

[0116] The present invention also relates to a recombinant polynucleotide, which contains a polynucleotide portion of a plant stress-regulated gene operatively linked to a heterologous nucleotide sequence. As used herein, the term "polynucleotide portion of plant stress-regulated sequence" means a contiguous nucleotide sequence of the plant stress-regulated gene that provides a function. The portion can be any portion of the sequence, particularly a coding sequence, or a sequence encoding a peptide portion of the stress-regulated polypeptide; the stress-regulated regulatory element; a sequence useful as an antisense molecule or triplexing agent; or a sequence useful for disrupting (knocking-out) an endogenous plant stress-regulated gene.

[0117] A heterologous nucleotide sequence is a nucleotide sequence that is not normally part of the plant stress-regulated gene from which the polynucleotide portion of the plant stress-regulated gene-component of the recombinant polynucleotide is obtained; or, if it is a part of the plant stress-regulated gene from which the polynucleotide portion is obtained, it is an orientation other than it would normally be in, for example, is an antisense sequence, or comprises at least partially discontinuous as compared to the genomic structure, for example, a single exon operatively linked to the regulatory element. In general, where the polynucleotide portion of the plant stress-regulated gene comprises the coding sequence in a recombinant polynucleotide of the invention, the heterologous nucleotide sequence will function as a regulatory element. The regulatory element can be any heterologous regulatory element, including, for example, a constitutively active regulatory element, an inducible regulatory element, or a tissue specific or phase specific regulatory element, as disclosed above. Conversely, where the polynucleotide portion of the plant stressregulated polynucleotide comprises the stress-regulated regulatory element of a recombinant polynucleotide of the invention, the heterologous nucleotide sequence generally will be a nucleotide sequence that can be transcribed and, if desired, translated. Where the heterologous nucleotide sequence is expressed from a plant stress-regulated regulatory element, it generally confers a desirable phenotype to a plant cell containing the recombinant polynucleotide, or provides a means to identify a plant cell containing the recombinant polynucleotide. It should be recognized that a "desirable" phenotype can be one that decreases the ability of a plant cell to compete where the plant cell, or a plant containing the cell, is an undesired plant cell. Thus, a heterologous nucleotide sequence can allow a plant to grow, for example, under conditions in which it would not normally be able to grow.

[0118] A heterologous nucleotide sequence can be, or encode, a selectable marker. As used herein, the term "selectable marker" is used herein to refer to a molecule that, when present or expressed in a plant cell, provides a means to identify a plant cell containing the marker. As such, a selectable marker can provide a means for screening a population of plants, or plant cells, to identify those having the marker. A

selectable marker also can confer a selective advantage to the plant cell, or a plant containing the cell. The selective advantage can be, for example, the ability to grow in the presence of a negative selective agent such as an antibiotic or herbicide, compared to the growth of plant cells that do not contain the selectable marker. The selective advantage also can be due, for example, to an enhanced or novel capacity to utilize an added compound as a nutrient, growth factor or energy source. A selectable advantage can be conferred, for example, by a single polynucleotide, or its expression product, or to a combination of polynucleotides whose expression in a plant cell gives the cell with a positive selective advantage, a negative selective advantage, or both.

Examples of selectable markers include those that confer antimetabolite [0119] resistance, for example, dihydrofolate reductase, which confers resistance to methotrexate (Reiss, Plant Physiol, (Life Sci. Adv.) 13:143-149, 1994); neomycin phosphotransferase, which confers resistance to the aminoglycosides neomycin, kanamycin and paromycin (Herrera-Estrella, EMBO J. 2:987-995, 1983) and hygro, which confers resistance to hygromycin (Marsh, Gene 32:481-485, 1984), trpB, which allows cells to utilize indole in place of tryptophan; hisD, which allows cells to utilize histinol in place of histidine (Hartman, Proc. Natl. Acad. Sci., USA 85:8047, 1988); mannose-6-phosphate isomerase which allows cells to utilize mannose (WO 94/20627); ornithine decarboxylase, which confers resistance to the ornithine decarboxylase inhibitor, 2-(difluoromethyl)-DL-ornithine (DFMO; McConlogue, 1987. In: Current Communications in Molecular Biology, Cold Spring Harbor Laboratory ed.); and deaminase from Aspergillus terreus, which confers resistance to Blasticidin S (Tamura, Biosci, Biotechnol, Biochem, 59:2336-2338, 1995). Additional selectable markers include those that confer herbicide resistance, for example, phosphinothricin acetyltransferase gene, which confers resistance to phosphinothricin (White et al., Nucl. Acids Res. 18:1062, 1990; Spencer et al., Theor. Appl. Genet. 79:625-631, 1990), a mutant EPSPV-synthase, which confers glyphosate resistance (Hinchee et al., Bio/Technology 91:915-922, 1998), a mutant acetolactate synthase, which confers imidazolione or sulfonylurea resistance (Lee et al., EMBO J. 7:1241-1248, 1988), a mutant psbA, which confers resistance to atrazine (Smeda et

al., <u>Plant Physiol.</u> 103:911-917, 1993), or a mutant protoporphyrinogen oxidase (see U.S. Pat. No. 5,767,373), or other markers conferring resistance to an herbicide such as glufosinate. In addition, markers that facilitate identification of a plant cell containing the polynucleotide encoding the marker include, for example, luciferase (Giacomin, <u>Plant Sci.</u> 116:59-72, 1996; Scikantha, <u>J. Bacteriol.</u> 178:121, 1996), green fluorescent protein (Gerdes, <u>FEBS Lett.</u> 389:44-47, 1996) or fl-glucuronidase (Jefferson, <u>EMBO J.</u> 6:3901-3907, 1997), and numerous others as disclosed herein or otherwise known in the art. Such markers also can be used as reporter molecules.

[0120] A heterologous nucleotide sequence can encode an antisense molecule, particularly an antisense molecule specific for a nucleotide sequence of a plant stressregulated gene, for example, the gene from which the regulatory component of the recombinant polynucleotide is derived. Such a recombinant polynucleotide can be useful for reducing the expression of a plant stress-regulated polypeptide in response to a stress condition because the antisense molecule, like the polypeptide, only will be induced upon exposure to the stress. A heterologous nucleotide sequence also can be, or can encode, a ribozyme or a triplexing agent. In addition to being useful as heterologous nucleotide sequences, such molecules also can be used directly in a method of the invention, for example, to modulate the responsiveness of a plant cell to a stress condition. Thus, an antisense molecule, ribozyme, or triplexing agent can be contacted directly with a target cell and, upon uptake by the cell, can effect their antisense, ribozyme or triplexing activity; or can be encoded by a heterologous nucleotide sequence that is expressed in a plant cell from a plant stress-regulated regulatory element, whereupon it can effect its activity.

[0121] An antisense polynucleotide, ribozyme or triplexing agent is complementary to a target sequence, which can be a DNA or RNA sequence, for example, messenger RNA, and can be a coding sequence, a nucleotide sequence comprising an intron-exon junction, a regulatory sequence such as a Shine-Delgarno-like sequence, or the like. The degree of complementarity is such that the polynucleotide, for example, an antisense polynucleotide, can interact specifically

with the target sequence in a cell. Depending on the total length of the antisense or other polynucleotide, one or a few mismatches with respect to the target sequence can be tolerated without losing the specificity of the polynucleotide for its target sequence. Thus, few if any mismatches would be tolerated in an antisense molecule consisting, for example, of twenty nucleotides, whereas several mismatches will not affect the hybridization efficiency of an antisense molecule that is complementary, for example, to the full length of a target mRNA encoding a cellular polypeptide. The number of mismatches that can be tolerated can be estimated, for example, using well known formulas for determining hybridization kinetics (see Sambrook et al., "Molecular Cloning; A Laboratory Manual" 2nd Edition (Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY; 1989)) or can be determined empirically using methods as disclosed herein or otherwise known in the art, particularly by determining that the presence of the antisense polynucleotide, ribozyme, or triplexing agent in a cell decreases the level of the target sequence or the expression of a polypeptide encoded by the target sequence in the cell.

[0122] A nucleotide sequence useful as an antisense molecule, a ribozyme or a triplexing agent can inhibit translation or cleave a polynucleotide encoded by plant stress-regulated gene, thereby modulating the responsiveness of a plant cell to a stress condition. An antisense molecule, for example, can bind to an mRNA to form a double stranded molecule that cannot be translated in a cell. Antisense oligonucleotides of at least about 15 to 25 nucleotides are preferred since they are easily synthesized and can hybridize specifically with a target sequence, although longer antisense molecules can be expressed from a recombinant polynucleotide introduced into the target cell. Specific nucleotide sequences useful as antisense molecules can be identified using well known methods, for example, gene walking methods (see, for example, Seimiya et al., J. Biol. Chem. 272:4631-4636 (1997), which is incorporated herein by reference). Where the antisense molecule is contacted directly with a target cell, it can be operatively associated with a chemically reactive group such as iron-linked EDTA, which cleaves a target RNA at the site of

hybridization. A triplexing agent, in comparison, can stall transcription (Maher et al., Antisense Res. Devel. 1:227 (1991); Helene. Anticancer Drug Design 6:569 (1991)).

[0123] A plant stress-regulated regulatory element can be included in an expression cassette. As used herein, the term "expression cassette" refers to a nucleotide sequence that can direct expression of an operatively linked polynucleotide. Thus, a plant stress-regulated regulatory element can constitute an expression cassette, or component thereof. An expression cassette is particularly useful for directing expression of a nucleotide sequence, which can be an endogenous nucleotide sequence or a heterologous nucleotide sequence, in a cell, particularly a plant cell. If desired, an expression cassette also can contain additional regulatory elements, for example, nucleotide sequences required for proper translation of a polynucleotide sequence into a polypeptide. In general, an expression cassette can be introduced into a plant cell such that the plant cell, a plant resulting from the plant cell, seeds obtained from such a plant, or plants produced from such seeds are resistant to a stress condition.

[0124] Additional regulatory sequences as disclosed above or other desirable sequences such as selectable markers or the like can be incorporated into an expression cassette containing a plant stress-regulated regulatory element (see, for example, WO 99/47552). Examples of suitable markers include dihydrofolate reductase (DHFR) or neomycin resistance for eukaryotic cells and tetracycline or ampicillin resistance for E. coli. Selection markers in plants include bleomycin, gentamycin, glyphosate, hygromycin, kanamycin, methotrexate, phleomycin, phosphinotricin, spectinomycin, streptomycin, sulfonamide and sulfonylureas resistance (see, for example, Maliga et al., Methods in Plant Molecular Biology, Cold Spring Harbor Laboratory Press, 1995, page 39). The selection marker can have its own promoter or its expression can be driven by the promoter operably linked to the sequence of interest. Additional sequences such as intron sequences (e.g. from Adh1 or bronze1) or viral leader sequences (e.g. from TMV, MCMV and AIVIV), all of which can enhance expression, can be included in the cassette. In addition, where it is

desirable to target expression of a nucleotide sequence operatively linked to the stressregulated regulatory element, a sequence encoding a cellular localization motif can be
included in the cassette, for example, such that an encoded transcript or translation
product is translocated to and localizes in the cytosol, nucleus, a chloroplast, or
another subcellular organelle. Examples of useful transit peptides and transit peptide
sequences can be found in Von Heijne et al., Plant Mol. Biol. Rep. 9: 104, 1991;
Clark et al., J. Biol. Chem. 264:17544, 1989; della Cioppa et al., Plant Physiol.
84:965, 1987; Romer et al., Biochem. Biophys. Res. Comm. 196:1414, 1993; Shah et
al., Science 233:478, 1986; Archer et al., J. Bioenerg Biomemb. 22:789, 1990;
Scandalios, Prog. Clin. Biol. Res. 344:515, 1990; Weisbeek et al., J. Cell Sci. Suppl.
11:199, 1989; Bruce, Trends Cell Biol. 10:440, 2000. The present invention can
utilize native or heterologous transit peptides. The encoding sequence for a transit
peptide can include all or a portion of the encoding sequence for a particular transit
peptide, and may also contain portions of the mature protein encoding sequence
associated with a particular transit peptide.

A polynucleotide portion of a plant stress-regulated plant gene, or an [0125] expression cassette, can be introduced into a cell as a naked DNA molecule, can be incorporated in a matrix such as a liposome or a particle such as a viral particle, or can be incorporated into a vector. Such vectors can be cloning or expression vectors, but other uses are within the scope of the present invention. A cloning vector is a selfreplicating DNA molecule that serves to transfer a DNA segment into a host cell. The three most common types of cloning vectors are bacterial plasmids, phages, and other viruses. An expression vector is a cloning vector designed so that a coding sequence inserted at a particular site will be transcribed and translated into a protein. Incorporation of the polynucleotide into a vector can facilitate manipulation of the polynucleotide, or introduction of the polynucleotide into a plant cell. A vector can be derived from a plasmid or a viral vector such as a T-DNA vector (Horsch et al., Science 227:1229-1231, 1985, which is incorporated herein by reference). If desired, the vector can comprise components of a plant transposable element, for example, a Ds transposon (Bancroft and Dean, Genetics 134:1221-1229, 1993, which is

incorporated herein by reference) or an Spm transposon (Aarts et al., Mol. Gen. Genet. 247:555-564, 1995, which is incorporated herein by reference).

In addition to containing the polynucleotide portion of a plant stressregulated gene, a vector can contain various nucleotide sequences that facilitate, for example, rescue of the vector from a transformed plant cell; passage of the vector in a host cell, which can be a plant, animal, bacterial, or insect host cell; or expression of an encoding nucleotide sequence in the vector, including all or a portion of a rescued coding region. As such, the vector can contain any of a number of additional transcription and translation elements, including constitutive and inducible promoters, enhancers, and the like (see, for example, Bitter et al., Meth. Enzymol. 153:516-544, 1987). For example, a vector can contain elements useful for passage, growth or expression in a bacterial system, including a bacterial origin of replication; a promoter, which can be an inducible promoter; and the like. In comparison, a vector that can be passaged in a mammalian host cell system can have a promoter such as a metallothionein promoter, which has characteristics of both a constitutive promoter and an inducible promoter, or a viral promoter such as a retrovirus long terminal repeat, an adenovirus late promoter, or the like. A vector also can contain one or more restriction endonuclease recognition and cleavage sites, including, for example, a polylinker sequence, to facilitate rescue of a nucleotide sequence operably linked to the polynucleotide portion.

[0127] The present invention also relates to a method of using a polynucleotide portion of a plant stress-regulated gene to confer a selective advantage on a plant cell. Such a method can be performed by introducing, for example, a plant stress-regulated regulatory element into a plant cell, wherein, upon exposure of the plant cell to a stress condition to which the regulatory element is responsive, a nucleotide sequence operatively linked to the regulatory element is expressed, thereby conferring a selective advantage to plant cell. The operatively linked nucleotide sequence can be a heterologous nucleotide sequence, which can be operatively linked to the regulatory element prior to introduction of the regulatory sequence into the plant cell; or can be

an endogenous nucleotide sequence into which the regulatory element was targeted by a method such as homologous recombination. The selective advantage conferred by the operatively linked nucleotide sequence can be such that the plant is better able to tolerate the stress condition; or can be any other selective advantage.

As used herein, the term "selective advantage" refers to the ability of a particular organism to better propagate, develop, grow, survive, or otherwise tolerate a condition as compared to a corresponding reference organism that does not contain a plant-stress regulated polynucleotide portion of the present invention. In one embodiment, a selective advantage is exemplified by the ability of a desired plant, plant cell, or the like, that contains an introduced plant stress-regulated regulatory element, to grow better than an undesired plant, plant cell, or the like, that does not contain the introduced regulatory element. For example, a recombinant polynucleotide comprising a plant stress-regulated regulatory element operatively linked to a heterologous nucleotide sequence encoding an enzyme that inactivates an herbicide can be introduced in a desired plant. Upon exposure of a mixed population of plants comprising the desired plants, which contain the recombinant polynucleotide, and one or more other populations of undesired plants, which lack the recombinant polynucleotide, to a stress condition that induces expression of the regulatory element and to the herbicide, the desired plants will have a greater likelihood of surviving exposure to the toxin and, therefore, a selective advantage over the undesired plants.

[0129] In another embodiment, a selective advantage is exemplified by the ability of a desired plant, plant cell, or the like, to better propagate, develop, grow, survive, or otherwise tolerate a condition as compared to an undesired plant, plant cell, or the like, that contains an introduced plant stress-regulated regulatory element. For example, a recombinant polynucleotide comprising a plant stress-regulated regulatory element operatively linked to a plant cell toxin can be introduced into cells of an undesirable plant present in a mixed population of desired and undesired plants, for example, food crops and weeds, respectively, then the plants can be exposed to stress

conditions that induce expression from the plant stress-regulated regulatory element. whereby expression of the plant cell toxin results in inhibition of growth or death of the undesired plants, thereby providing a selective advantage to the desired plants. which no longer have to compete with the undesired plants for nutrients, light, or the like. In another example, a plant stress-regulated regulatory element operatively linked to a plant cell toxin can be introduced into cells of plants used as a nurse crop. Nurse crops, also called cover or companion crops, are planted in combination with plants of interest to provide, among other things, shade and soil stability during establishment of the desired plants. Once the desired plants have become established, the presence of the nurse crop may no longer be desirable. Exposure to conditions inducing expression of the gene linked to the plant stress-regulated regulatory element allows elimination of the nurse crop. Alternatively nurse crops can be made less tolerate to abiotic stress by the inhibition of any of the stress-regulated sequences disclosed herein. Inhibition can be accomplished by any of the method described herein. Upon exposure of the nurse crop to the stress, the decreased ability of the nurse crop to respond to the stress will result in elimination of the nurse crop, leaving only the desired plants.

[0130] The invention also provides a means of producing a transgenic plant, which comprises plant cells that exhibit altered responsiveness to a stress condition. As such, the present invention further provides a transgenic plant, or plant cells or tissues derived therefrom, which are genetically modified to respond to stress differently than a corresponding wild-type plant or plant not containing constructs of the present invention would respond. As used herein, the term "responsiveness to a stress condition" refers to the ability of a plant to express a plant stress-regulated gene upon exposure to the stress condition. A transgenic plant cell contains a polypeptide portion of a plant stress-regulated gene, or a mutant form thereof, for example, a knock-out mutant. A knock-out mutant form of a plant stress-regulated gene can contain, for example, a mutation such that a STOP codon is introduced into the reading frame of the translated portion of the gene such that expression of a functional stress-regulated polypeptide is prevented; or a mutation in the stress-regulated

regulatory element such that inducibility of the element in response to a stress condition is inhibited. Such transgenic plants of the invention can display any of various idiotypic modifications is response to an abiotic stress, including altered tolerance to the stress condition, as well as increased or decreased plant growth, root growth, yield, or the like, as compared to the corresponding wild-type plant.

[0131] The term "plant" is used broadly herein to include any plant at any stage of development, or to part of a plant, including a plant cutting, a plant cell, a plant cell culture, a plant organ, a plant seed, and a plantlet. A plant cell is the structural and physiological unit of the plant, comprising a protoplast and a cell wall. A plant cell can be in the form of an isolated single cell or a cultured cell, or can be part of higher organized unit, for example, a plant tissue, plant organ, or plant. Thus, a plant cell can be a protoplast, a gamete producing cell, or a cell or collection of cells that can regenerate into a whole plant. As such, a seed, which comprises multiple plant cells and is capable of regenerating into a whole plant, is considered plant cell for purposes of this disclosure. A plant tissue or plant organ can be a seed, protoplast, callus, or any other groups of plant cells that is organized into a structural or functional unit. Particularly useful parts of a plant include harvestable parts and parts useful for propagation of progeny plants. A harvestable part of a plant can be any useful part of a plant, for example, flowers, pollen, seedlings, tubers, leaves, stems, fruit, seeds, roots, and the like. A part of a plant useful for propagation includes, for example, seeds, fruits, cuttings, seedlings, tubers, rootstocks, and the like.

[0132] A transgenic plant can be regenerated from a transformed plant cell. As used herein, the term "regenerate" means growing a whole plant from a plant cell; a group of plant cells; a protoplast; a seed; or a piece of a plant such as a callus or tissue. Regeneration from protoplasts varies from species to species of plants. For example, a suspension of protoplasts can be made and, in certain species, embryo formation can be induced from the protoplast suspension, to the stage of ripening and germination. The culture media generally contains various components necessary for growth and regeneration, including, for example, hormones such as auxins and

cytokinins; and amino acids such as glutamic acid and proline, depending on the particular plant species. Efficient regeneration will depend, in part, on the medium, the genotype, and the history of the culture. If these variables are controlled, however, regeneration is reproducible.

[0133] Regeneration can occur from plant callus, explants, organs or plant parts. Transformation can be performed in the context of organ or plant part regeneration. (see Meth-Enzymol. Vol. 118; Klee et al. <a href="Ann. Rev. Plant Physiol, 38:467, 1987, which is incorporated herein by reference). Utilizing the leaf disk-transformation-regeneration method, for example, disks are cultured on selective media, followed by shoot formation in about two to four weeks (see Horsch et al., supra, 1985). Shoots that develop are excised from calli and transplanted to appropriate root-inducing selective medium. Rooted plantlets are transplanted to soil as soon as possible after roots appear. The plantlets can be repotted as required, until reaching maturity.

[0134] In vegetatively propagated crops, the mature transgenic plants are propagated utilizing cuttings or tissue culture techniques to produce multiple identical plants. Selection of desirable transgenotes is made and new varieties are obtained and propagated vegetatively for commercial use. In seed propagated crops, the mature transgenic plants can be self crossed to produce a homozygous inbred plant. The resulting inbred plant produces seeds that contain the introduced plant stress-induced regulatory element, and can be grown to produce plants that express a polynucleotide or polypeptide in response to a stress condition that induces expression from the regulatory element. As such, the invention further provides seeds produced by a transgenic plant obtained by a method of the invention.

[0135] In addition, transgenic plants comprising different recombinant sequences can be crossbred, thereby providing a means to obtain transgenic plants containing two or more different transgenes, each of which contributes a desirable characteristic to the plant. Methods for breeding plants and selecting for crossbred plants having desirable characteristics or other characteristics of interest are well known in the art. [0136] A method of the invention can be performed by introducing a polynucleotide portion of a plant stress-regulated gene into the plant. As used herein, the term "introducing" means transferring a polynucleotide into a plant cell. A polynucleotide can be introduced into a cell by a variety of methods well known to those of ordinary skill in the art. For example, the polynucleotide can be introduced into a plant cell using a direct gene transfer method such as electroporation or microprojectile mediated transformation, or using Agrobacterium mediated transformation. Non-limiting examples of methods for the introduction of polynucleotides into plants are provided in greater detail herein. As used herein, the term "transformed" refers to a plant cell containing an exogenously introduced polynucleotide portion of a plant stress-regulated gene that is or can be rendered active in a plant cell, or to a plant comprising a plant cell containing such a polynucleotide.

[0137] It should be recognized that one or more polynucleotides, which are the same or different can be introduced into a plant, thereby providing a means to obtain a genetically modified plant containing multiple copies of a single transgenic sequence, or containing two or more different transgenic sequences, either or both of which can be present in multiple copies. Such transgenic plants can be produced, for example, by simply selecting plants having multiple copies of a single type of transgenic sequence; by cotransfecting plant cells with two or more populations of different transgenic sequences and identifying those containing the two or more different transgenic sequences; or by crossbreeding transgenic plants, each of which contains one or more desired transgenic sequences, and identifying those progeny having the desired sequences.

[0138] Methods for introducing a polynucleotide into a plant cell to obtain a transformed plant also include direct gene transfer (see European Patent A 164 575), injection, electroporation, biolistic methods such as particle bombardment, pollenmediated transformation, plant RNA virus-mediated transformation, liposomemediated transformation, transformation using wounded or enzyme-degraded immature embryos, or wounded or enzyme-degraded embryogenic callus, and the like. Transformation methods using Agrobacterium tumefactens tumor inducing (Ti) plasmids or root-inducing (Ri) plasmids, or plant virus vectors are well known in the art (see, for example, WO 99/47552; Weissbach & Weissbach, "Methods for Plant Molecular Biology" (Academic Press, NY 1988), section VIII, pages 421-463; Grierson and Corey, "Plant Molecular Biology" 2d Ed. (Blackie, London 1988), Chapters 7-9, each of which is incorporated herein by reference; Horsch et al., supra, 1985). The wild-type form of Agrobacterium, for example, contains a Ti plasmid, which directs production of tumorigenic crown gall growth on host plants. Transfer of the tumor inducing T-DNA region of the Ti plasmid to a plant genome requires the Ti plasmid-encoded virulence genes as well as T-DNA borders, which are a set of direct DNA repeats that delineate the region to be transferred. An Agrobacterium based vector is a modified form of a Ti plasmid, in which the tumor inducing functions are replaced by a nucleotide sequence of interest that is to be introduced into the plant host.

[0139] Methods of using Agrobacterium mediated transformation include cocultivation of Agrobacterium with cultured isolated protoplasts; transformation of plant cells or tissues with Agrobacterium; and transformation of seeds, apices or meristems with Agrobacterium. In addition, in planta transformation by Agrobacterium can be performed using vacuum infiltration of a suspension of Agrobacterium cells (Bechtold et al., C.R. Acad. Sci. Paris 316:1194, 1993, which is incorporated herein by reference).

[0140] Agrobacterium mediated transformation can employ cointegrate vectors or binary vector systems, in which the components of the Ti plasmid are divided between a helper vector, which resides permanently in the Agrobacterium host and carries the virulence genes, and a shuttle vector, which contains the gene of interest bounded by T-DNA sequences. Binary vectors are well known in the art (see, for example, De Framond, BioTechnology 1:262, 1983; Hoekema et al., Nature 303:179, 1983, each of which is incorporated herein by reference) and are commercially

available (Clontech; Palo Alto CA). For transformation, Agrobacterium can be cocultured, for example, with plant cells or wounded tissue such as leaf tissue, root explants, hypocotyledons, stem pieces or tubers (see, for example, Glick and Thompson, "Methods in Plant Molecular Biology and Biotechnology" (Boca Raton FL, CRC Press 1993), which is incorporated herein by reference). Wounded cells within the plant tissue that have been infected by Agrobacterium can develop organs de novo when cultured under the appropriate conditions; the resulting transgenic shoots eventually give rise to transgenic plants, which contain an exogenous polynucleotide portion of a plant stress-regulated gene.

[0141] Agrobacterium mediated transformation has been used to produce a variety of transgenic plants, including, for example, transgenic cruciferous plants such as Arabidopsis, mustard, rapeseed and flax; transgenic leguminous plants such as alfalfa, pea, soybean, trefoil and white clover; and transgenic solanaceous plants such as eggplant, petunia, potato, tobacco and tomato (see, for example, Wang et al., "Transformation of Plants and Soil Microorganisms" (Cambridge, University Press 1995), which is incorporated herein by reference). In addition, Agrobacterium mediated transformation can be used to introduce an exogenous polynucleotide sequence, for example, a plant stress-regulated regulatory element into apple, aspen, belladonna, black currant, carrot, celery, cotton, cucumber, grape, horseradish, lettuce, morning glory, muskmelon, neem, poplar, strawberry, sugar beet, sunflower, walnut, asparagus, rice and other plants (see, for example, Glick and Thompson, supra, 1993; Hiei et al., Plant J. 6:271-282, 1994; Shimamoto, Science 270:1772-1773, 1995).

[0142] Suitable strains of Agrobacterium tumefaciens and vectors as well as transformation of Agrobacteria and appropriate growth and selection media are well known in the art (GV3101, pMK90RK), Koncz, Mol. Gen. Genet. 204:383-396, 1986; (C58C1, pGV3850kan), Deblaere, Nucl. Acid Res. 13:4777, 1985; Bevan, Nucl. Acid Res. 12:8711, 1984; Koncz, Proc. Natl. Acad. Sci. USA 86:8467-8471, 1986; Koncz, Plant Mol. Biol. 20:963-976, 1992; Koncz, Specialized vectors for gene tagging and expression studies. In: Plant Molecular Biology Manual Vol. 2, Gelvin and

Schilperoort (Eds.), Dordrecht, The Netherlands: Kluwer Academic Publ. (1994), 1-22; European Patent A-1 20 516; Hoekema: The Binary Plant Vector System, Offsetdrukkerij Kanters B. V., Alblasserdam (1985), Chapter V; Fraley, <u>Crit. Rev. Plant. Sci.</u>, 4:1-46; An. EMBO J. 4:277-287, 1985).

[0143] Where a polynucleotide portion of a plant stress-regulated gene is contained in vector, the vector can contain functional elements, for example "left border" and "right border" sequences of the T-DNA of Agrobacterium, which allow for stable integration into a plant genome. Furthermore, methods and vectors that permit the generation of marker-free transgenic plants, for example, where a selectable marker gene is lost at a certain stage of plant development or plant breeding, are known, and include, for example, methods of co-transformation (Lyznik, Plant Mol. Biol. 13:151-161, 1989; Peng, Plant Mol. Biol. 27:91-104, 1995), or methods that utilize enzymes capable of promoting homologous recombination in plants (see, e.g., W097/08331; Bayley, Plant Mol. Biol. 18:353-361, 1992; Lloyd, Mol. Gen. Genet. 242:653-657, 1994; Maeser, Mol. Gen. Genet. 230:170-176, 1991; Onouchi, Nucl. Acids Res. 19:6373-6378, 1991; see, also, Sambrook et al., supra, 1989).

10144] A direct gene transfer method such as electroporation also can be used to introduce a polynucleotide portion of a plant stress-regulated gene into a cell such as a plant cell. For example, plant protoplasts can be electroporated in the presence of the regulatory element, which can be in a vector (Fromm et al., Proc. Natl. Acad. Sci., USA 82:5824, 1985, which is incorporated herein by reference). Electrical impulses of high field strength reversibly permeabilize membranes allowing the introduction of the nucleic acid. Electroporated plant protoplasts reform the cell wall, divide and form a plant callus. Microinjection can be performed as described in Potrykus and Spangenberg (eds.), Gene Transfer To Plants (Springer Verlag, Berlin, NY 1995). A transformed plant cell containing the introduced polynucleotide can be identified by detecting a phenotype due to the introduced polynucleotide, for example, increased or decreased tolerance to a stress condition.

[0145] Microprojectile mediated transformation also can be used to introduce a polynucleotide into a plant cell (Klein et al., Nature 327:70-73, 1987, which is incorporated herein by reference). This method utilizes microprojectiles such as gold or tungsten, which are coated with the desired nucleic acid molecule by precipitation with calcium chloride, spermidine or polyethylene glycol. The microprojectile particles are accelerated at high speed into a plant tissue using a device such as the BIOLISTIC PD-1000 (BioRad: Hercules CA).

Microprojectile mediated delivery ("particle bombardment") is especially [0146] useful to transform plant cells that are difficult to transform or regenerate using other methods. Methods for the transformation using biolistic methods are well known (Wan, Plant Physiol. 104:37-48, 1984; Vasil, Bio/Technology 11:1553-1558, 1993; Christou, Trends in Plant Science 1:423-431, 1996). Microprojectile mediated transformation has been used, for example, to generate a variety of transgenic plant species, including cotton, tobacco, corn, hybrid poplar and papaya (see Glick and Thompson, supra, 1993). Important cereal crops such as wheat, oat, barley, sorghum and rice also have been transformed using microprojectile mediated delivery (Duan et al., Nature Biotech, 14:494-498, 1996; Shimamoto, Curr. Opin. Biotech, 5:158-162, 1994). A rapid transformation regeneration system for the production of transgenic plants such as a system that produces transgenic wheat in two to three months (see European Patent No. EP 0709462A2, which is incorporated herein by reference) also can be useful for producing a transgenic plant using a method of the invention, thus allowing more rapid identification of gene functions. The transformation of most dicotyledonous plants is possible with the methods described above. Transformation of monocotyledonous plants also can be transformed using, for example, biolistic methods as described above, protoplast transformation, electroporation of partially permeabilized cells, introduction of DNA using glass fibers, Agrobacterium mediated transformation, and the like.

[0147] Plastid transformation also can be used to introduce a polynucleotide portion of a plant stress-regulated gene into a plant cell (U.S. Patent Nos. 5.451.513,

5,545,817, and 5,545,818; WO 95/16783; McBride et al., Proc. Natl. Acad. Sci., USA 91:7301-7305, 1994). Chloroplast transformation involves introducing regions of cloned plastid DNA flanking a desired nucleotide sequence, for example, a selectable marker together with polynucleotide of interest into a suitable target tissue, using, for example, a biolistic or protoplast transformation method (e.g., calcium chloride or PEG mediated transformation). One to 1.5 kb flanking regions ("targeting sequences") facilitate homologous recombination with the plastid genome, and allow the replacement or modification of specific regions of the plastome. Using this method, point mutations in the chloroplast 16S rRNA and rps12 genes, which confer resistance to spectinomycin and streptomycin, can be utilized as selectable markers for transformation (Syab et al., Proc. Natl. Acad. Sci., USA 87:8526-8530, 1990; Staub and Maliga, Plant Cell 4:39-45, 1992), resulted in stable homopiasmic transformants; at a frequency of approximately one per 100 bombardments of target leaves. The presence of cloning sites between these markers allowed creation of a plastid targeting vector for introduction of foreign genes (Staub and Maliga, EMBO J. 12:601-606, 1993). Substantial increases in transformation frequency are obtained by replacement of the recessive rRNA or r-protein antibiotic resistance genes with a dominant selectable marker, the bacterial aadA gene encoding the spectinomycindetoxifying enzyme aminoglycoside-3'-adenyltransf erase (Svab and Maliga, Proc. Natl. Acad. Sci., USA 90:913-917, 1993). Approximately 15 to 20 cell division cycles following transformation are generally required to reach a homoplastidic state. Plastid expression, in which genes are inserted by homologous recombination into all of the several thousand copies of the circular plastid genome present in each plant cell, takes advantage of the enormous copy number advantage over nuclear-expressed genes to permit expression levels that can readily exceed 10% of the total soluble plant protein.

[0148] Plants suitable to treatment according to a method of the invention can be monocots or dicots and include, but are not limited to, corn (*Zea mays*), *Brassica* sp. (e.g., *B. napus*, *B. rapa*, *B. juncea*), particularly those *Brassica* species useful as sources of seed oil, alfalfa (*Medicago sativa*), rice (*Oryza sativa*), rye (*Secale*

cereale), sorghum (Sorghum bicolor, Sorghum vulgare), millet (e.g., pearl millet (Pennisetum glaucum), proso millet (Panicum miliaceum), foxtail millet (Setaria italica), finger millet (Eleusine coracana)), sunflower (Helianthus annuus), safflower (Carthamus tinctorius), wheat (Triticum aestivum), soybean (Glycine max), tobacco (Nicotiana tabacum), potato (Solanum tuberosum), peanuts (Arachis hypogaea), cotton (Gossypium barbadense, Gossypium hirsutum), sweet potato (Ipomoea batatus), cassava (Manihot esculenta), coffee (Cofea spp.), coconut (Cocos nucifera), pineapple (Ananas comosus), citrus trees (Citrus spp.), cocoa (Theobroma cacao), tea (Camellia sinensis), banana (Musa spp.), avocado (Persea ultilane), fig (Ficus casica), guava (Psidium guajava), mango (Mangifera indica), olive (Olea europaea), papaya (Carica papaya), cashew (Anacardium occidentale), macadamia (Macadamia integrifolia), almond (Prunus amvgdalus), sugar beets (Beta vulgaris), sugarcane (Saccharum spp.), oats, duckweed (Lemna), barley, tomatoes (Lycopersicon esculentum), lettuce (e.g., Lactuca sativa), green beans (Phaseolus vulgaris), lima beans (Phaseolus limensis), peas (Lathyrus spp.), and members of the genus Cucumis such as cucumber (C. sativus), cantaloupe (C. cantalupensis), and musk melon (C. melo).

[0149] Ornamentals such as azalea (Rhododendron spp.), hydrangea (Macrophylla hydrangea), hibiscus (Hibiscus rosasanensis), roses (Rosa spp.), tulips (Tulipa spp.), daffodils (Narcissus spp.), petunias (Petunia hybrida), carnation (Dianthus caryophyllus), poinsettia (Euphorbia pulcherrima), and chrysanthemum are also included. Additional ornamentals within the scope of the invention include impatiens, Begonia, Pelargonium, Viola, Cyclamen, Verbena, Vinca, Tagetes, Primula, Saint Paulia, Agertum, Amaranthus, Antihirrhinum, Aquilegia, Cineraria, Clover, Cosmo, Cowpea, Dahlia, Datura, Delphinium, Gerbera, Gladiolus, Gloxinia, Hippeastrum, Mesembryanthemum, Salpiglossos, and Zinnia.

[0150] Conifers that may be employed in practicing the present invention include, for example, pines such as loblolly pine (*Pinus taeda*), slash pine (*Pinus elliotii*), ponderosa pine (*Pinus ponderosa*), lodgepole pine (*Pinus contorta*), and Monterey

pine (Pinus radiata), Douglas-fir (Pseudotsuga menziesii); Western hemlock (Tsuga ultilane); Sitka spruce (Picea glauca); redwood (Sequoia sempervirens); true firs such as silver fir (Abies amabilis) and balsam fir (Abies balsamea); and cedars such as Western red cedar (Thuja plicata) and Alaska yellow-cedar (Chamaecyparis nootkatensis).

[0151] Leguminous plants which may be used in the practice of the present invention include beans and peas. Beans include guar, locust bean, fenugreek, soybean, garden beans, cowpea, mungbean, lima bean, fava bean, lentils, chickpea, etc. Legumes include, but are not limited to, Arachis, e.g., peanuts, Vicia, e.g., crown vetch, hairy vetch, adzuki bean, mung bean, and chickpea, Lupinus, e.g., lupine, trifolium, Phaseolus, e.g., common bean and lima bean, Pisum, e.g., field bean, Melilotus, e.g., clover, Medicago, e.g., alfalfa, Lotus, e.g., trefoil, lens, e.g., lentil, and false indigo. Preferred forage and turf grass for use in the methods of the invention include alfalfa, orchard grass, tall fescue, perennial ryegrass, creeping bent grass, and redtop.

[0152] Other plants within the scope of the invention include Acacia, aneth, artichoke, arugula, blackberry, canola, cilantro, clementines, escarole, eucalyptus, fennel, grapefruit, honey dew, jicama, kiwifruit, lemon, lime, mushroom, nut, okra, orange, parsley, persimmon, plantain, pomegranate, poplar, radiata pine, radicchio, Southern pine, sweetgum, tangerine, triticale, vine, yams, apple, pear, quince, cherry, apricot, melon, hemp, buckwheat, grape, raspberry, chenopodium, blueberry, nectarine, peach, plum, strawberry, watermelon, eggplant, pepper, cauliflower, Brassica, e.g., broccoli, cabbage, ultilan sprouts, onion, carrot, leek, beet, broad bean, celery, radish, pumpkin, endive, gourd, garlic, snapbean, spinach, squash, turnip, ultilane, chicory, groundnut and zucchini.

[0153] Angiosperms are divided into two broad classes based on the number of cotyledons, which are seed leaves that generally store or absorb food; a monocotyledonous angiosperm has a single cotyledon, and a dicotyledonous angiosperm has two cotyledons. Angiosperms produce a variety of useful products including materials such as lumber, rubber, and paper; fibers such as cotton and linen; herbs and medicines such as quinine and vinblastine; ornamental flowers such as roses and orchids; and foodstuffs such as grains, oils, fruits and vegetables.

[0154] Angiosperms encompass a variety of flowering plants, including, for example, cereal plants, leguminous plants, oilseed plants, hardwood trees, fruit-bearing plants and ornamental flowers, which general classes are not necessarily exclusive. Cereal plants, which produce an edible grain cereal, include, for example, corn, rice, wheat, barley, oat, rye, orchardgrass, guinea grass, sorghum and turfgrass. Leguminous plants include members of the pea family (Fabaceae) and produce a characteristic fruit known as a legume. Examples of leguminous plants include, for example, soybean, pea, chickpea, moth bean, broad bean, kidney bean, lima bean, lentil, cowpea, dry bean, and peanut, as well as alfalfa, birdsfoot trefoil, clover and sainfoin. Oilseed plants, which have seeds that are useful as a source of oil, include soybean, sunflower, rapeseed (canola) and cottonseed.

[0155] Angiosperms also include hardwood trees, which are perennial woody plants that generally have a single stem (trunk). Examples of such trees include alder, ash, aspen, basswood (linden), beech, birch, cherry, cottonwood, elm, eucalyptus, hickory, locust, maple, oak, persimmon, poplar, sycamore, walnut, sequoia, and willow. Trees are useful, for example, as a source of pulp, paper, structural material and firel

[0156] Angiosperms are fruit-bearing plants that produce a mature, ripened ovary, which generally contains seeds. A fruit can be suitable for human or animal consumption or for collection of seeds to propagate the species. For example, hops are a member of the mulberry family that are prized for their flavoring in malt liquor. Fruit-bearing angiosperms also include grape, orange, lemon, grapefruit, avocado, date, peach, cherry, olive, plum, coconut, apple and pear trees and blackberry, blueberry, raspberry, strawberry, pineapple, tomato, cucumber and eggplant plants.

An ornamental flower is an angiosperm cultivated for its decorative flower. Examples of commercially important ornamental flowers include rose, orchid, lily, tulip and chrysanthemum, snapdragon, camellia, carnation and petunia plants. The skilled artisan will recognize that the methods of the invention can be practiced using these or other angiosperms, as desired, as well as gymnosperms, which do not produce seeds in a fruit.

[0157] A method of producing a transgenic plant can be performed by introducing a polynucleotide portion of plant stress-regulated gene into a plant cell genome, whereby the polynucleotide portion of the plant stress-regulated gene modulates a response of the plant cell to a stress condition, thereby producing a transgenic plant, which comprises plant cells that exhibit altered responsiveness to the stress condition. In one embodiment, the polynucleotide portion of the plant stress-regulated gene encodes a stress-regulated polypeptide or functional peptide portion thereof, wherein expression of the stress-regulated polypeptide or functional peptide portion thereof either increases the stress tolerance of the transgenic plant, or decreases the stress tolerance of the transgenic plant. The polynucleotide portion of the plant stress-regulated gene encoding the stress-regulated polypeptide or functional peptide portion thereof can be operatively linked to a heterologous promoter.

[0158] In another embodiment, the polynucleotide portion of the plant stress-regulated gene comprises a stress-regulated regulatory element. The stress-regulated regulatory element can integrate into the plant cell genome in a site-specific manner, whereupon it can be operatively linked to an endogenous nucleotide sequence, which can be expressed in response to a stress condition specific for the regulatory element; or can be a mutant regulatory element, which is not responsive to the stress condition, whereby upon integrating into the plant cell genome, the mutant regulatory element disrupts an endogenous stress-regulated regulatory element of a plant stress-regulated gene, thereby altering the responsiveness of the plant stress-regulated gene to the stress condition. Accordingly, the invention also provides genetically modified plants, including transgenic plants, produced by such a method, and a plant cell

obtained from such genetically modified plant, wherein said plant cell exhibits altered responsiveness to the stress condition; a seed produced by a transgenic plant; and a cDNA library prepared from a transgenic plant.

[0159] Also provided is a method of modulating the responsiveness of a plant cell to a stress condition. Such a method can be performed, for example, by introducing a polynucleotide portion of a plant stress-regulated gene into the plant cell, thereby modulating the responsiveness of the plant cell to a stress condition. As disclosed herein, the responsiveness of the plant cell can be increased or decreased upon exposure to the stress condition, and the altered responsiveness can result in increased or decreased tolerance of the plant cell to a stress condition. The polynucleotide portion of the plant stress-regulated gene can, but need not, be integrated into the genome of the plant cell, thereby modulating the responsiveness of the plant cell to the stress condition. Accordingly, the invention also provide a genetically modified plant, including a transgenic plant, which contains an introduced polynucleotide portion of a plant stress-regulated gene, as well as plant cells, tissues, and the like, which exhibit modulated responsiveness to a stress condition.

[0160] The polynucleotide portion of the plant stress-regulated gene can encode a stress-regulated polypeptide or functional peptide portion thereof, which can be operatively linked to a heterologous promoter. As used herein, reference to a "functional peptide portion of a plant stress-regulated polypeptide" means a contiguous amino acid sequence of the polypeptide that has an activity of the full length polypeptide, or that has an antagonist activity with respect to the full length polypeptide, or that presents an epitope unique to the polypeptide. Thus, by expressing a functional peptide portion of a plant stress-regulated polypeptide in a plant cell, the peptide can act as an agonist or an antagonist of the polypeptide, thereby modulating the responsiveness of the plant cell to a stress condition.

[0161] A polynucleotide portion of the plant stress-regulated nucleotide sequence also can contain a mutation, whereby upon integrating into the plant cell genome, the

polynucleotide disrupts (knocks-out) an endogenous plant stress-regulated nucleotide sequence, thereby modulating the responsiveness of said plant cell to the stress condition. Depending on whether the knocked-out gene encodes an adaptive or a maladaptive stress-regulated polypeptide, the responsiveness of the plant will be modulated accordingly. Thus, a method of the invention provides a means of producing a transgenic plant having a knock-out phenotype of a plant stress-regulated nucleotide sequence.

[0162] Alternatively, the responsiveness of a plant or plant cell to a stress condition can be modulated by use of a suppressor construct containing dominant negative mutation for any of the stress-regulated sequences described herein. Expression of a suppressor construct containing a dominant mutant mutation generates a mutant transcript that, when coexpressed with the wild-type transcript inhibits the action of the wild-type transcript. Methods for the design and use of dominant negative constructs are well known (see, for example, in Herskowitz, Nature 329:219-222, 1987; Lagna and Hemmati-Brivanlou, Curr. Topics Devel. Biol. 36:75-98, 1998).

[0163] The polynucleotide portion of the plant stress-regulated gene also can comprise a stress-regulated regulatory element, which can be operatively linked to a heterologous nucleotide sequence, which, upon expression from the regulatory element in response to a stress condition, modulates the responsiveness of the plant cell to the stress condition. Such a heterologous nucleotide sequence can encode, for example, a stress-inducible transcription factor such as DREB1A, which, upon exposure to the stress condition, is expressed such that it can amplify the stress response (see Kasuga et al., supra, 1999). The heterologous nucleotide sequence also can encode a polynucleotide that is specific for a plant stress-regulated gene, for example, an antisense molecule, a ribozyme, and a triplexing agent, either of which, upon expression in the plant cell, reduces or inhibits expression of a stress-regulated polypeptide encoded by the gene, thereby modulating the responsiveness of the plant cell to a stress condition, for example, an abnormal level of cold, osmotic pressure,

and salinity. As used herein, the term "abnormal," when used in reference to a condition such as temperature, osmotic pressure, salinity, or any other condition that can be a stress condition, means that the condition varies sufficiently from a range generally considered optimum for growth of a plant that the condition results in an induction of a stress response in a plant. Methods of determining whether a stress response has been induced in a plant are disclosed herein or otherwise known in the art

[0164] A plant stress-regulated regulatory element can be operatively linked to a heterologous polynucleotide sequence, such that the regulatory element can be introduced into a plant genome in a site-specific matter by homologous recombination. For example, a mutant plant stress-regulated regulatory element for a maladaptive stress-induced polypeptide can be transformed into a plant genome in a site specific manner by in vivo mutagenesis, using a hybrid RNA-DNA oligonucleotide ("chimeroplast" (TIBTECH 15:441-447, 1997; W0 95/15972; Kren, Hepatology 25:1462-1468, 1997; Cole-Strauss, Science 273:1386-1389, 1996, each of which is incorporated herein by reference). Part of the DNA component of the RNA-DNA oligonucleotide is homologous to a nucleotide sequence comprising the regulatory element of the maladaptive gene, but includes a mutation or contains a heterologous region which is surrounded by the homologous regions. By means of base pairing of the homologous regions of the RNA-DNA oligonucleotide and of the endogenous nucleic acid molecule, followed by a homologous recombination the mutation contained in the DNA component of the RNA-DNA oligonucleotide or the heterologous region can be transferred to the plant genome, resulting in a "mutant" gene that, for example, is not induced in response to a stress and, therefore, does not confer the maladaptive phenotype. Such a method similarly can be used to knock-out the activity of a stress-regulated gene, for example, in an undesirable plant. Such a method can provide the advantage that a desirable wild-type plant need not compete with the undesirable plant, for example, for light, nutrients, or the like.

[0165] A method of modulating the responsiveness of a plant cell to a stress condition also can be performed by introducing a mutation in the chromosomal copy of a plant stress-regulated gene, for example, in the stress-regulated regulatory element, by transforming a cell with a chimeric oligonucleotide composed of a contiguous stretch of RNA and DNA residues in a duplex conformation with double hairpin caps on the ends. An additional feature of the oligonucleotide is the presence of 2'-0- methylation at the RNA residues. The RNA/DNA sequence is designed to align with the sequence of a chromosomal copy of the target regulatory element and to contain the desired nucleotide change (see U.S. Pat. No. 5,501,967, which is incorporated herein by reference).

[0166] A plant stress-regulated regulatory element also can be operatively linked to a heterologous polynucleotide such that, upon expression from the regulatory element in the plant cell, confers a desirable phenotype on the plant cell. For example, the heterologous polynucleotide can encode an aptamer, which can bind to a stress-induced polypeptide. Aptamers are nucleic acid molecules that are selected based on their ability to bind to and inhibit the activity of a protein or metabolite. Antamers can be obtained by the SELEX (Systematic Evolution of Ligands by Exponential Enrichment) method (see U.S. Pat. No. 5,270,163), wherein a candidate mixture of single stranded nucleic acids having regions of randomized sequence is contacted with a target, and those nucleic acids having a specific affinity to the target are partitioned from the remainder of the candidate mixture, and amplified to yield a ligand enriched mixture. After several iterations a nucleic acid molecule (aptamer) having optimal affinity for the target is obtained. For example, such a nucleic acid molecule can be operatively linked to a plant stress-regulated regulatory element and introduced into a plant. Where the aptamer is selected for binding to a polypeptide that normally is expressed from the regulatory element and is involved in an adaptive response of the plant to a stress, the recombinant molecule comprising the aptamer can be useful for inhibiting the activity of the stress-regulated polypeptide, thereby decreasing the tolerance of the plant to the stress condition.

101671 The invention provides a genetically modified plant, which can be a transgenic plant, that is tolerant or resistant to a stress condition. As used herein, the term "tolerant" or "resistant," when used in reference to a stress condition of a plant, means that the particular plant, when exposed to a stress condition, shows less of an effect, or no effect, in response to the condition as compared to a corresponding reference plant (naturally occurring wild-type plant or a plant not containing a construct of the present invention). As a consequence, a plant encompassed within the present invention grows better under more widely varying conditions, has higher yields and/or produces more seeds. Thus, a transgenic plant produced according to a method of the invention can demonstrate protection (as compared to a corresponding reference plant) from a delay to complete inhibition of alteration in cellular metabolism, or reduced cell growth or cell death caused by the stress. Preferably, the transgenic plant is capable of substantially normal growth under environmental conditions where the corresponding reference plant shows reduced growth, metabolism or viability, or increased male or female sterility.

The determination that a plant modified according to a method of the [0168] invention has increased resistance to a stress-inducing condition can be made by comparing the treated plant with a control (reference) plant using well known methods. For example, a plant having increased tolerance to saline stress can be identified by growing the plant on a medium such as soil, which contains a higher content of salt in the order of at least about 10% compared to a medium the corresponding reference plant is capable of growing on. Advantageously, a plant treated according to a method of the invention can grow on a medium or soil containing at least about 50%, or more than about 75%, particularly at least about more than 100%, and preferably more than about 200% salt than the medium or soil on which a corresponding reference plant can grow. In particular, such a treated plant can grow on medium or soil containing at least 40 mM, generally at least 100 mM, particularly at least 200 mM, and preferably at least 300 mM salt, including, for example, a water soluble inorganic salt such as sodium sulfate, magnesium sulfate, calcium sulfate, sodium chloride, magnesium chloride, calcium chloride, potassium

chloride, or the like; salts of agricultural fertilizers, and salts associated with alkaline or acid soil conditions; particularly NaCl.

[0169] In another embodiment, the invention provides a plant that is less tolerant or less resistant to a stress condition as compared to a corresponding reference plant. As used herein, the term "less tolerant" or "less resistant," when used in reference to a stress condition of a plant, means that the particular plant, when exposed to a stress condition, shows an alteration in response to the condition as compared to a corresponding reference plant. As a consequence, such a plant, which generally is an undesirable plant species, is less likely to grow when exposed to a stress condition than an untreated plant.

[0170] The present invention also relates to a method of expressing a heterologous nucleotide sequence in a plant cell. Such a method can be performed, for example, by introducing into the plant cell a plant stress-regulated regulatory element operatively linked to the heterologous nucleotide sequence, whereby, upon exposure of the plant cell to stress condition, the heterologous nucleotide sequence is expressed in the plant cell. The heterologous nucleotide sequence can encode a selectable marker, or preferably, a polypeptide that confers a desirable trait upon the plant cell, for example, a polypeptide that improves the nutritional value, digestibility or ornamental value of the plant cell, or a plant comprising the plant cell. Accordingly, the invention provides a transgenic plant that, in response to a stress condition, can produce a heterologous polypeptide from a plant stress-regulated regulatory element. Such transgenic plants can provide the advantage that, when grown in a cold environment for example, expression of the heterologous polypeptide from a plant cold-regulated regulatory element can result in increased nutritional value of the plant.

[0171] The present invention further relates to a method of modulating the activity of a biological pathway in a plant cell, wherein the pathway involves a stressregulated polypeptide. As used herein, reference to a pathway that "involves" a stress-regulated polypeptide means that the polypeptide is required for normal function of the pathway. For example, plant stress-regulated polypeptides as disclosed herein include those acting as kinases or as transcription factors, which are well known to be involved in signal transduction pathways. As such, a method of the invention provides a means to modulate biological pathways involving plant stress-regulated polypeptides, for example, by altering the expression of the polypeptides in response to a stress condition. Thus, a method of the invention can be performed, for example, by introducing a polynucleotide portion of a plant stress-regulated gene into the plant cell, thereby modulating the activity of the biological pathway.

[0172] A method of the invention can be performed with respect to a pathway involving any of the stress-regulated polypeptides as encoded by a polynucleotide of SEQ ID NOS:1-2703, including for example, a stress-regulated transcription factor, an enzyme, including a kinase, a channel protein (see, for example, Tables 29-31; see, also, Table 1). Pathways in which the disclosed stress-regulated stress factors are involved can be identified, for example, by searching the Munich Information Center for Protein Sequences (MIPS) *Arabidopsis thaliana* database (MATDB), which is at http://www.mips.biochem.mpg.de/proj/thal/.

[0173] The present invention also relates to a method of identifying a polynucleotide that modulates a stress response in a plant cell. Such a method can be performed, for example, by contacting an array of probes representative of a plant cell genome and nucleic acid molecules expressed in plant cell exposed to the stress; detecting a nucleic acid molecule that is expressed at a level different from a level of expression in the absence of the stress; introducing the nucleic acid molecule that is expressed differently into a plant cell; and detecting a modulated response of the plant cell containing the introduced nucleic acid molecule to a stress, thereby identifying a polynucleotide that modulates a stress response in a plant cell. The contacting is under conditions that allow for selective hybridization of a nucleic acid molecule with probe having sufficient complementarity, for example, under stringent hybridization conditions.

[0174] As used herein, the term "array of probes representative of a plant cell genome" means an organized group of oligonucleotide probes that are linked to a solid support, for example, a microchip or a glass slide, wherein the probes can hybridize specifically and selectively to nucleic acid molecules expressed in a plant cell. Such an array is exemplified herein by a GeneChip® Arabidopsis Genome Array (Affymetrix; see Example 1). In general, an array of probes that is "representative" of a plant genome will identify at least about 30% or the expressed nucleic acid molecules in a plant cell, generally at least about 50% or 70%, particularly at least about 80% or 90%, and preferably will identify all of the expressed nucleic acid molecules. It should be recognized that the greater the representation, the more likely all nucleotide sequences of cluster of stress-regulated genes will be identified.

[0175] A method of the invention is exemplified in Example 1, wherein clusters of Arabidopsis genes induced to cold, to increased salinity, to increased osmotic pressure, and to a combination of the above three stress conditions were identified. Based on the present disclosure, the artisan readily can obtain nucleic acid samples for Arabidopsis plants exposed to other stress conditions, or combinations of stress conditions, and identify clusters of genes induced in response to the stress conditions. Similarly, the method is readily adaptable to identifying clusters of stress-regulated genes expressed in other plant species, particularly commercially valuable plant species, where a substantial amount of information is known regarding the genome.

[0176] The clusters of genes identified herein include those clusters of genes that are induced or repressed in response to a combination of stress conditions, but not to any of the stress conditions alone; and clusters of genes that are induced or repressed in response to a selected stress condition, but not to other stress conditions tested. Furthermore, clusters of genes that respond to a stress condition in a temporally regulated manner are also included, such as gene clusters that are induced early (for example, within about 3 hours), late (for example, after about 8 to 24 hours), or continuously in a stress response. In addition, the genes within a cluster are represented by a variety of cellular proteins, including transcription factors, enzymes

such as kinases, channel proteins, and the like (see Tables 1 and 29-31). Thus, the present invention further characterizes nucleotide sequences that previously were known to encode cellular peptides by classifying them within clusters of stress-regulated genes.

[0177] The present invention additionally relates to a method of identifying a stress condition to which a plant cell was exposed. Such a method can be performed, for example, by contacting nucleic acid molecules expressed in the plant cell and an array of probes representative of the plant cell genome; and detecting a profile of expressed nucleic acid molecules characteristic of a stress response, thereby identifying the stress condition to which the plant cell was exposed. The contacting generally is under conditions that allow for selective hybridization of a nucleic acid molecule with probe having sufficient complementarity, for example, under stringent hybridization conditions. The profile can be characteristic of exposure to a single stress condition, for example, an abnormal level of cold, osmotic pressure, or salinity (Tables 3-14), or can be characteristic of exposure to more than one stress condition (Tables 15-26, for example, cold, increased osmotic pressure and increased salinity (see Tables 24-26).

[0178] The method can be practiced using at least one nucleic acid probe and can identify one or combination of stress conditions by detecting altered expression of one or a plurality of polynucleotides representative of plant stress-regulated genes. As used herein, the term "at least one" includes one, two, three or more, for example, five, ten, twenty, fifty or more polynucleotides, nucleic acid probes, and the like. The term "plurality" is used herein to mean two or more, for example, three, four, five or more, including ten, twenty, fifty or more polynucleotides, nucleic acid probes, and the like.

[0179] In a method of the invention, nucleic acid samples from the plant cells to be collected can be contacted with an array, then the profile can be compared with known expression profiles prepared from nucleic acid samples of plants exposed to a

known stress condition or combination of stress conditions. By creating a panel of such profiles, representative of various stress conditions, an unknown stress condition to which a plant was exposed can be identified simply by comparing the unknown profile with the known profiles and determining which known profile that matches the unknown profile. Preferably, the comparison is automated. Such a method can be useful, for example, to identify a cause of damage to a crop, where the condition causing the stress is not known or gradually increases over time. For example, accumulation in soils over time of salts from irrigation water can result in gradually decreasing crop yields. Because the accumulation is gradual, the cause of the decreased yield may not be readily apparent. Using the present methods, it is possible to evaluate the stress to which the plants are exposed, thus revealing the cause of the decreased yields.

[0180] The present invention, therefore includes a computer readable medium containing executable instructions form receiving expression data for sequences substantially similar to any of those disclosed herein and comparing expression data from a test plant to a reference plant that has been exposed to an abiotic stress. Also provided is a computer-readable medium containing sequence data for sequences substantially similar to any of the sequences described herein, or the complements thereof, and a module for comparing such sequences to other nucleic acid sequences.

[0181] Also provided are plants and plant cells comprising plant stress-regulatory elements of the present invention operably linked to a nucleotide sequence encoding a detectable signal. Such plants can be used as diagnostic or "sentinel" plants to provide early warning that nearby plants are being stressed so that appropriate actions can be taken. In one embodiment, the signal is one that alters the appearance of the plant. For example, an osmotic stress regulatory element of the present invention can be operably linked to a nucleotide sequence encoding a fluorescent protein such as green fluorescent protein. When subjected to osmotic stress, the expression of the green fluorescent protein in the sentinel plant provides a visible signal so that appropriate actions can be taken to remove or alleviate the stress. The use of

fluorescent proteins in plants is well known (see, for example, in Leffel et al., BioTechniques 23:912, 1997).

[0182] The invention further relates to a method of identifying an agent that modulates the activity of a stress-regulated regulatory element of a plant. As used herein, the term "modulate the activity," when used in reference to a plant stress-regulated regulatory element, means that expression of a polynucleotide from the regulatory element is increased or decreased. In particular, expression can be increased or decreased with respect to the basal activity of the promoter, i.e., the level of expression, if any, in the absence of a stress condition that normally induces expression from the regulatory element; or can be increased or decreased with respect to the level of expression in the presence of the inducing stress condition. As such, an agent can act as a mimic of a stress condition, or can act to modulate the response to a stress condition.

[0183] Such a method can be performed, for example, by contacting the regulatory element with an agent suspected of having the ability to modulate the activity of the regulatory element, and detecting a change in the activity of the regulatory element. In one embodiment, the regulatory element can be operatively linked to a heterologous polynucleotide encoding a reporter molecule, and an agent that modulates the activity of the stress-regulated regulatory element can be identified by detecting a change in expression of the reporter molecule due to contacting the regulatory element with the agent. Such a method can be performed *in vitro* in a plant cell-free system, or in a plant cell in culture or in a plant *in situ*.

[0184] A method of the invention also can be performed by contacting the agent is contacted with a genetically modified cell or a transgenic plant containing an introduced plant stress-regulated regulatory element, and an agent that modulates the activity of the regulatory element is identified by detecting a phenotypic change in the modified cell or transgenic plant.

A method of the invention can be performed in the presence or absence of [0185] the stress condition to which the particularly regulatory element is responsive. As such, the method can identify an agent that modulates the activity of plant stressregulated promoter in response to the stress, for example, an agent that can enhance the stress response or can reduce the stress response. In particular, a method of the invention can identify an agent that selectively activates the stress-regulated regulatory elements of a cluster of plant stress-regulated genes, but does not affect the activity of other stress-regulated regulatory genes. As such, the method provides a means to identify an agent that acts as a stress mimic. Such agents can be particularly useful to prepare a plant to an expected stress condition. For example, a agent that acts as a cold mimic can be applied to a field of plants prior to the arrival of an expected cold front. Thus, the cold stress response can be induced prior to the actual cold weather, thereby providing the plants with the protection of the stress response, without the plants suffering from any initial damage due to the cold. Similarly, an osmotic pressure mimic can be applied to a crop of plants prior a field being flooded by a rising river.

[0186] In one embodiment, the present invention provides a method for markerassisted selection. Marker-assisted selection involves the selection of plants having
desirable phenotypes based on the presence of particular nucleotide sequences
("markers"). The use of markers allows plants to be selected early in development,
often before the phenotype would normally be manifest. Because it allows for early
selection, marker-assisted selection decreases the amount of time need for selection
and thus allows more rapid genetic progress.

[0187] Briefly, marker-assisted selection involves obtaining nucleic acid from a plant to be selected. The nucleic acid obtained is then probed with probes that selectively hybridize under stringent, preferably highly stringent, conditions to a nucleotide sequence or sequences associated with the desired phenotype. In one embodiment, the probes hybridize to any of the stress-responsive genes or regulatory regions disclosed herein, for example, any one of SEQ ID NOS:1-2703. The presence

of any hybridization products formed is detected and plants are then selected on the presence or absence of the hybridization products.

[0188] The following examples are intended to illustrate but not limit the invention

EXAMPLE 1

PROFILING OF PLANT STRESS-REGULATED GENES

[0189] This example demonstrates that clusters of stress-regulated genes can be identified in plant cells exposed to various stress conditions, either alone or in combination

[0190] A GeneChip® Arabidopsis Genome Array (Affymetrix, Santa Clara, CA) was used to identify clusters of genes that were coordinately induced in response to various stress conditions. The GeneChip® Arabidopsis Genome Array contains probes synthesized in situ and is designed to measure temporal and spatial gene expression of approximately 8700 genes in greater than 100 EST clusters. The sequences used to develop the array were obtained from GenBank (http://www.ncbi.nlm.nih.gov/) in collaboration with Torrey Mesa Research Institute (San Diego, CA), formerly known as Novartis Agriculture Discovery Institute. Eighty percent of the nucleotide sequences represented on the array are predicted coding sequences from genomic BAC entries; twenty percent are high quality cDNA sequences. The array also contains over 100 EST clusters that share homology with the predicted coding sequences from BAC clones (see, for example, world wide web at address (url) "affymetrix.com/products/Arabidopsis_content.html".

[0191] The Affymetrix GeneChip® array was used to define nucleotide sequences/ pathways affected by various abiotic stresses and to define which are uniquely regulated by one stress and those that respond to multiple stress, and to identify candidate nucleotide sequences for screening for insertional mutants. Of the approximately 8,700 nucleotide sequences represented on the Affymetrix GeneChip® array, 2862 nucleotide sequences showed at least a 2-fold change in expression in at least one sample, relative to no-treatment controls. Of those 2,862 nucleotide sequences 1,335 were regulated only by cold stress, 166 were regulated only mannitol stress and 209 were regulated only by saline stress. Furthermore, of the 2,862 nucleotide sequences 123 nucleotide sequences were regulated by salt and mannitol stress, 293 were regulated by mannitol and cold stress, 274 were regulated by cold and saline stress and 462 were regulated by cold, mannitol and salt. Of the 2,862 nucleotide sequences, 771 passed the higher stringency of showing at least a 2-fold change in expression in at least 2 samples, relative to control. And, 508 of the 771 nucleotide sequences were found in an in-house collection of insertion mutants.

[0192] The following describes in more detail how the experiments were done. Transcriptional profiling was performed by hybridizing fluorescence labeled cRNA with the oligonucleotides probes on the chip, washing, and scanning. Each gene is represented on the chip by about sixteen oligonucleotides (25-mers). Expression level is related to fluorescence intensity. Starting material contained 1 to 10 μ g total RNA; detection specificity was about 1:10 6 ; approximately a 2-fold change was detectable, with less than 2% false positive; the dynamic range was approximately 500x. Nucleotide sequences having up to 70% to 80% identity could be discriminated using this system.

[0193] Seven day old axenic *Arabidopsis* seedlings were transferred to Magenta boxes with rafts floating on MS medium. Three weeks later (28 day old seedlings), stresses were applied as follows: Control - no treatment; Cold - Magenta box placed in ice; Mannitol - medium + 200 mM mannitol; Salt - medium + 100 mM NaCl. Tissue samples were collected at 3 hours and 27 hours into the stress, roots and aerial portions were harvested, RNA was purified, and the samples were analyzed using the GeneChip® Arabidopsis Genome Array (Affymetrix, Santa Clara, CA) following the manufacturer's protocol.

[0194] Raw fluorescence values as generated by Affymetrix software were processed as follows: the values were brought into Microsoft Excel and values of

25 or less were set to 25 (an empirically determined baseline, Zhu and Wang, Plant Physiol. 124:1472-1476; 2000). The values from the stressed samples were then converted to fold change relative to control by dividing the values from the stressed samples by the values from the no-treatment control samples. Expression patterns that were altered at least 2-fold with respect to the control were selected. This method gave very robust results and resulted in a larger number of nucleotide sequences called as stress-regulated than previous methods had permitted.

[0195] Based on the profiles obtained following hybridization of nucleic acid molecules obtained from plant cells exposed to various stress conditions to the probes in the microarray, clusters of nucleotide sequences that were altered in response to the stress conditions were identified (see Tables 3-6, cold responsive; Tables 7-10, salt (saline) responsive; Tables 11 to 14, mannitol (osmotic) responsive; Tables 15-17, cold and mannitol responsive; Tables 18-20, 6 salt and cold responsive; Tables 21-23, salt and mannitol responsive; Tables 24-26, cold, salt and mannitol responsive. Examples of plant gene sequences that varied in expression at least two-fold in response to a combination of cold, saline and osmotic stress in root cells and leaf cells are shown in Tables 27 and 28, respectively. In addition, examples of plant gene sequences that encode transcription factors (Table 29), phosphatases (Table 30), and kinases (Table 31) and that varied at least two-fold in response to a combination of cold, saline and osmotic stress are provided.

[0196] Affymetrix ID numbers and corresponding SEQ ID NOS: for the respective Arabidopsis nucleotide sequences are provided Tables 3-26, and can be used to determine SEQ ID NOS: for the sequences shown by Affymetrix ID number in Tables 27-31. The Affymetrix ID number refers to a particular nucleotide sequence on the GeneChip® Arabidopsis Genome Array. In some cases, a particular plant stress-regulated gene sequence hybridized to more than one nucleotide sequence on the GeneChip® Arabidopsis Genome Array (see, for example, Table 3, where SEQ ID NO:36 is shown to have hybridized to the 12187_AT and 15920_I_AT nucleotide sequences on the GeneChip®). In addition, it should be recognized that the disclosed

sequences are not limited to coding sequences but, in some cases, include 5' untranslated sequences (see Table 2) or a longest coding region. As such, while the sequences set forth as SEQ ID NOS:1-2073 generally start with an ATG codon, in most cases each comprises a longer nucleotide sequence, including a regulatory region (see Table 2).

The results disclosed herein demonstrate that several polynucleotides, some 101971 of which were known to function as transcription factors, enzymes, and structural proteins, also are involved in the response of a plant cell to stress. The identification of the clusters of stress-regulated genes as disclosed herein provides a means to identify stress-regulated regulatory elements present in Arabidopsis thaliana nucleotide sequences, including consensus regulatory elements. It should be recognized, however that the regulatory elements of the plant genes comprising a sequence as set forth in SEQ ID NOS:156, 229, 233, 558, 573, 606, 625, 635, 787, and 813, which previously have been described as cold regulated genes, are not encompassed within the stress-regulated gene regulatory element of the invention, and the regulatory elements of the plant genes comprising the nucleotide sequences set forth as SEQ ID NOS:1263, 1386, 1391, 1405, 1445, 1484, 1589, 1609, 1634, 1726, 1866, 1918, and 1928, which previously have been identified as genes that are responsive to a single stress condition such as cold or saline stress, are not encompassed within the plant stress-regulated gene regulatory elements of the invention to the extent that they confer stress-regulated expression only with respect to the known single stress. Furthermore, the identification of the Arabidopsis stressregulated genes provides a means to identify the corresponding homologs and orthologs in other plants, including commercially valuable food crops such as wheat, rice, soy, and barley, and ornamental plants. BLASTN and BLASTP searches to identify such sequences revealed the polynucleotide sequences set forth in Table 32, which is on the CD-R compact disc submitted herewith.

[0198] Although the invention has been described with reference to the above example, it will be understood that modifications and variations are encompassed within the spirit and scope of the invention. Accordingly, the invention is limited only by the claims, which follow Tables 1 to 31.

TABLE 1

SEQUENCE DESCRIPTIONS

Seq	Description	41	scarecrow-like 7 (SCL7)
ID		42	putative protein
1	unknown protein	43	No function assigned by TIGR
2	unknown protein	44	unknown protein
3	unknown protein	45	unknown protein
4	putative auxin-induced		
proteir	1	SEQ	Description
5	unknown protein	ID	
6	hypothetical protein	46	succinyl-CoA-ligase alpha subunit
7	putative protein	47	putative protein
8	unknown protein	48	CLV1 receptor kinase like protein
9	unknown protein	49	putative receptor-like protein
10	unknown protein		kinase
11	putative protein	50	putative squalene synthase
12	Thioredoxin - like protein	51	putative receptor protein kinase
13	putative RNA helicase	52	somatic embryogenesis receptor-
14	putative protein		like kinase, putative
15	putative protein	53	putative protein
16	RING zinc finger protein,	54	putative beta-glucosidase
	putative	55	multi-drug resistance protein
17	putative cyclin	56	receptor protein kinase (TMK1),
18	putative protein		putative
19	putative protein	57	putative receptor-like protein
20	unknown protein		kinase
21	putative protein	58	putative pectate lyase
22	putative protein	59	putative protein kinase
23	hypothetical protein	60	putative peroxidase
24	unknown protein	61	cytochrome P450-like protein
25	hypothetical protein	62	putative beta-amylase
26	unknown protein	63	monosaccharide transporter STP3
27	unknown protein	64	Lycopersicon esculentum
28	unknown protein		proteinase TMP, Pir2:T07617
29	unknown protein	65	putative receptor-like protein
30	putative protein		kinase
31	putative protein	66	G-box-binding factor 1
32	putative protein	67	amino acid carrier, putative
33	unknown protein	68	myb-related protein
34	putative ribonuclease III	69	No function assigned by TIGR
35	unknown protein	70	SNF1 like protein kinase
36	unknown protein	71	Cu/Zn superoxide dismutase-like
37	unknown protein		protein
38	unknown protein	72	putative protein kinase
39	unknown protein	73	small nuclear ribonucleoprotein
40	putative histidine kinase		U1A

		101	to the trade of the title of the
74	ras-like GTP-binding	101	dynein light chain like protein
proteir		102	chaperonin CPN10
75	oleoyl-[acyl-carrier-protein]	103	putative bHLH transcription factor
	hydrolase-like protein	104	putative glyoxysomal malate
76	putative heat shock		dehydrogenase precursor
	transcription factor	105	ATP-dependent RNA helicase,
77	putative protein		putative
78	membrane-bound small	106	chlorophyll synthetase
	GTP-binding - like protein	107	similar to epoxide hydrolases
79	putative protein (fragment)	108	putative protein
80	indole-3-acetate beta-	109	unknown protein
	glucosyltransferase like	110	hypothetical protein
	protein	111	putative membrane transporter
81	HD-zip transcription factor	112	putative tyrosyl-tRNA synthetase
	(athb-8)	113	ARGININE/SERINE-RICH
82	putative cAMP-dependent		SPLICING FACTOR RSP31
	protein kinase	114	putative oxidoreductase
83	glucuronosyl transferase-	115	unknown protein
	like protein	116	linker histone protein, putative
84	putative leucine-rich repeat	117	hypothetical protein
	disease resistance protein	118	putative protein
85	98b like protein	119	putative mitochondrial carrier
86	putative receptor-like		protein
	protein kinase	120	putative transcription factor
87	IAA-Ala hydrolase (IAR3)	121	MYB-related protein
88	putative AP2 domain	122	myb-related transcription factor,
	transcription factor		putative
89	putative expansin	123	unknown protein
90	putative Ap2 domain	124	unknown protein
prote		125	putative glycine-rich protein
91	expansin (At-EXP1)	126	No function assigned by TIGR
92	cytochrome P450 - like	127	unknown protein
prote		128	unknown protein
93	putative ATP-dependent	129	unknown protein
	RNA helicase A	130	unknown protein
94	unknown protein	131	putative membrane channel protein
95	predicted protein	132	putative protein
96	putative glucosyltransferase	133	unknown protein
97	unknown protein	134	gamma glutamyl hydrolase,
98	putative xyloglucan-		putative
	specific glucanase	135	40S ribosomal protein S5
99	cysteine synthase	136	DnaJ-like protein
100	clathrin assembly protein	137	40S ribosomal protein S26
	AP19 homolog	138	putative WRKY-type DNA binding
			protein

139	putative protein	161	putative photomorphogenesis
140	hypothetical protein		repressor protein
141	putative ubiquitin-	162	SNF1-like protein kinase (AKin11)
	conjugating enzyme	163	thioredoxin h
142	peptidylprolyl isomerase	164	thioredoxin
ROC1		165	Ca2+-dependent lipid-binding
143	glyceraldehyde-3-		protein, putative
	phosphate dehydrogenase C	166	putative auxin-induced protein
	subunit (GapC)	167	putative bZIP transcription factor
144	No function assigned by	168	hypothetical protein
TIGR		169	putative AVR9 elicitor response
145	putative protein		protein
146	putative thioredoxin	170	putative serine/threonine protein
147	thioredoxin h, putative		kinase
148	thioredoxin-like	171	bZIP transcription factor ATB2
149	allene oxide synthase	172	putative spliceosome associated
	(emb CAA73184.1)		protein
150	anthranilate synthase	173	3-hydroxyisobutyryl-coenzyme A
	component I-1 precursor		hydrolase - like protein
	(sp P32068)	174	putative protein
151	CELL DIVISION	175	putative Mutator-like transposase
	CONTROL PROTEIN 2	176	putative protein
	HOMOLOG A	177	unknown protein
152	protein kinase cdc2	178	putative protein
homo	log B	179	putative protein
153	ethylene responsive	180	putative galactinol synthase
	element binding factor 1	181	putative transcriptional regulator
	(frameshift!)	182	nuclear matrix constituent protein 1
154	ethylene responsive		(NMCP1)-like
	element binding factor 2	183	putative DNA-binding protein
	(ATERF2) (sp O80338)		RAV2
155	ethylene responsive	184	No function assigned by TIGR
	element binding factor 5	185	basic blue protein, 5' partial
	(ATERF5) (sp O80341)	186	unknown protein
156	glucose-6-phosphate	187	putative calcium-binding protein,
	dehydrogenase		calreticulin
157	photomorphogenesis	188	putative pyrophosphate-fructose-6-
	repressor (COP1)		phosphate 1-phosphotransferase
158	unknown protein	189	ribosomal protein L11, cytosolic
159	DNA (cytosine-5)-	190	putative dTDP-glucose 4-6-
	methyltransferase (DNA		dehydratase
	methyltransferase) (DNA	191	40S ribosomal protein S20-like
	metase) (sp P34881)		protein
160	PROLIFERA	192	60S ribosomal protein L24

193	coatomer-like protein,	223	putative SF16 protein {Helianthus
1,,,	epsilon subunit		annuus}
194	glycoprotein(EP1), putative	224	unknown protein
195	putative SPL1-related	225	thioredoxin
protein	1	226	trehalose-6-phosphate phosphatase
196	unknown protein		(AtTPPB)
197	putative transport protein	227	chlorophyll a/b-binding protein
	SEC61 beta-subunit	228	class IV chitinase (CHIV)
198	unknown protein	229	chalcone synthase (naringenin-
199	putative cytochrome P450		chalcone synthase) (testa 4 protein)
200	UTP-glucose		(sp P13114)
	glucosyltransferase - like	230	unknown protein
	protein	231	cinnamyl-alcohol dehydrogenase
201	60S ribosomal protein L23		ELI3-2
202	40S ribosomal protein S17	232	farnesyl-pyrophosphate synthetase
203	40S ribosomal protein S26		FPS2
204	protein translation factor	233	phospholipid hydroperoxide
	Sui1 homolog, putative		glutathione peroxidase
205	unknown protein	234	heat shock transcription factor
206	gamma glutamyl hydrolase,		HSF4
	putative	235	heat shock protein 101
207	dTDP-glucose 4,6-	236	17.6 kDa heat shock protein (AA
	dehydratase, putative		1-156)
208	extensin - like protein	237	heat shock protein 17.6A
209	unknown protein	238	heat-shock protein
210	protein phosphatase 2C -	239	HY5
	like protein	240	putative auxin-induced protein,
211	ubiquitin-like protein		IAA12
212	protein phosphatase 2C-like	241	early auxin-induced protein,
	protein		IAA19
213	unknown protein	242	auxin-inducible gene (IAA2)
214	putative RING zinc finger	243	putative protein
ankyr	in protein	244	putative choline kinase
215	unknown protein	245	thymidylate kinase - like protein
216	putative rubisco subunit	246	CTP synthase like protein
	binding-protein alpha	247	putative protein
	subunit	248	putative amidase
217	putative acetone-	249	4-alpha-glucanotransferase
	cyanohydrin lyase	250	hypothetical protein
218	putative isoamylase	251	similar to auxin-induced protein
219	putative protein	252	putative protein
220	HSP associated protein like	253	putative protein
221	60S ribosomal protein L39	254	putative protein
222	unknown protein	255	hyuC-like protein

256	putative tetracycline	287	unknown protein
	transporter protein	288	putative esterase D
257	similar to early nodulins	289	predicted protein of unknown
258	putative protein	function	on
259	putative peptidyl-prolyl cis-	290	unknown protein
	trans isomerase	291	putative indole-3-glycerol
260	unknown protein		phosphate synthase
261	unknown protein	292	isopentenyl
262	putative endochitinase		pyrophosphate:dimethyllallyl
263	putative ABC transporter		pyrophosphate isomerase
264	No function assigned by	293	kinase associated protein
TIGR			phosphatase
265	CONSTANS-like B-box	294	putative K+ channel, beta subunit
203	zinc finger protein	295	KNAT1 homeobox-like protein
266	unknown protein	296	PSI type II chlorophyll a/b-binding
267	unknown protein		protein, putative
268	putative mitochondrial	297	transcription factor
208	processing peptidase alpha	298	putative WD-40 repeat protein,
	subunit	2,0	MSI2
269	putative pre-mRNA	299	WD-40 repeat protein (MSI3)
20)	splicing factor	300	putative WD-40 repeat protein,
270	putative phosphatidylserine		MSI4
2,0	decarboxylase	301	unknown protein
271	unknown protein	302	hypothetical protein
272	unknown protein	303	putative protein
273	unknown protein	304	No function assigned by TIGR
274	putative casein kinase I	305	polyphosphoinositide binding
275	unknown protein		protein, putative
276	60S ribosomal protein	306	hypothetical protein
L23A		307	unknown protein
277	putative mitochondrial	308	chloroplast ribosomal L1 - like
	dicarboxylate carrier		protein
	protein	309	cold-regulated protein cor15b
278	enoyl-ACP reductase (enr-		precursor
A)	•,	310	cyanohydrin lyase like protein
279	putative isoamylase	311	putative replication protein A1
280	formamidase - like protein	312	putative protein
281	reticuline oxidase - like	313	possible apospory-associated like
prote	in		protein
282		314	DNA binding protein GT-1,
283	unknown protein		
	unknown protein putative transketolase	314	putative
	putative transketolase	315	
284	putative transketolase irsor		putative
	putative transketolase		putative AT-hook DNA-binding protein

318	enoyl-CoA hydratase like	348	putative farnesylated protein
	protein	349	unknown protein
319	berberine bridge enzyme -	350	water stress-induced protein,
	like protein		putative
320	putative sugar transporter	351	unknown protein
321	unknown protein	352	unknown protein
322	No function assigned by	353	PEROXISOMAL MEMBRANE
TIGR	9		PROTEIN PMP22
323	hypothetical protein	354	putative peroxisomal membrane
324	putative acidic ribosomal		carrier protein
	protein	355	putative protein
325	putative protein	356	unknown protein
326	unknown protein	357	putative protein
327	hypothetical protein	358	putative protein
328	putative protein	359	argininosuccinate synthase -like
329			protein
	dihydroxypolypreny	360	1-phosphatidylinositol-4,5-
	lbenzoate methyltransferase	bispho	osphate phosphodiesterase
330	unknown protein	361	putative JUN kinase activator
331	myb-related protein	protei	n
332	No function assigned by	362	putative 60S ribosomal protein L35
TIGR		363	nucleoid DNA-binding protein
333	putative protein		cnd41 - like protein
334	putative disease resistance	364	SigA binding protein
	response protein	365	hypothetical protein
335	hypothetical protein	366	putative protein kinase
336	No function assigned by	367	unknown protein
TIGR		368	regulatory protein NPR1-like;
337	starch branching enzyme II		transcription factor inhibitor I
338	No function assigned by		kappa B-like
TIGR		369	putative protein
339	putative enolase (2-	370	hypothetical protein
	phospho-D-glycerate	371	phosphoribosylanthranilate
	hydroylase)		isomerase
340	putative protein kinase	372	phosphoribosylanthranilate
341	HD-Zip protein, putative		isomerase
342	putative protein kinase	373	sterol glucosyltransferase, putative
343	phenylalanyl-trna	374	putative gigantea protein
	synthetase - like protein	375	putative MYB family transcription
344	putative aconitase		factor
345	NAM(no apical meristem)	376	hypothetical protein
	protein, putative	377	hypothetical protein
346	unknown protein	378	predicted protein
347	putative	379	cytochrome P450, putative
phosp	homannomutase		

	IA.	DEE I (COM	•9
380	putative Na+ dependent		chloroplast precursor (sp Q02166)
500	ileal bile acid transporter		phytochrome C (sp P14714)
381	unknown protein		putative phytochrome-associated
382	RING-H2 finger protein		protein 3
302	RHF1a		receptor serine/threonine kinase
383	putative protein	710	PR5K
384	unknown protein	419	Ran-binding protein (atranbp1a)
385		420	small Ras-like GTP-binding
	putative protein	420	protein (gb AAB58478.1)
386	putative auxin-regulated	421	sterol-C5-desaturase
205	protein		tryptophan synthase beta chain 1
387	hypothetical protein	422	
388	unknown protein	422	precursor (sp[P14671)
389	unknown protein	423	thioredoxin f2 (gb AAD35004.1)
390	putative protein	424	No function assigned by TIGR
391	putative protein	425	putative WRKY DNA-binding
392	unknown protein		protein
393	histone H1	426	putative protein
394	Argonaute (AGO1)-like	427	unknown protein
protein		428	unknown protein
395	unknown protein	429	14-3-3 protein homolog RCI1
396	putative protein with C-		(pir S47969)
	terminal RING finger	430	unknown protein
397	unknown protein	431	putative CCCH-type zinc finger
398	unknown protein	proteir	
399	unknown protein	432	PINHEAD (gb AAD40098.1);
400	unknown protein		tion initiation factor
401	unknown protein	433	plasma membrane proton ATPase
402	putative copper amine	(PMA)	
oxida	se	434	CHLOROPHYLL A-B BINDING
403	unknown protein		PROTEIN 4 PRECURSOR
404	unknown protein		homolog
405	unknown protein	435	membrane related protein CP5,
406	putative protein		putative
407	putative protein	436	ABC transporter (AtMRP2)
408	unknown protein	437	putative embryo-abundant protein
409	unknown protein	438	putative anthocyanidin-3-glucoside
410	putative protein		rhamnosyltransferase
411	putative protein	439	putative lipid transfer protein
412	unknown protein	440	unknown protein
413	serine/threonine kinase -	441	unknown protein
	like protein	442	galactinol synthase, putative
414	alcohol dehydrogenase,	443	putative protein
	putative	444	putative protein
415	anthranilate	445	SCARECROW-like protein
	phosphoribosyltransferase,	446	unknown protein

447	unknown protein	476	phosphoenolpyruvate carboxylase
448	unknown protein	477	(PPC) chlorophyll a/b-binding protein -
449	unknown protein	4//	like
450	asparaginetRNA ligase	470	AtAGP4
451	putative protein	478	
452	glutamate-1-semialdehyde	479	putative cryptochrome 2 apoprotein
	2,1-aminomutase 1	480	type 2 peroxiredoxin, putative
	precursor (GSA 1)	481	Atpm24.1 glutathione S transferase
	(glutamate-1-semialdehyde	482	delta tonoplast integral protein
	aminotransferase 1) (GSA-	400	(delta-TIP)
	AT 1) (sp P42799)	483	20S proteasome subunit (PAA2)
453	hypothetical protein	484	dormancy-associated protein,
454	putative serine protease-like	40.5	putative
	protein	485	putative cytidine deaminase
455	No function assigned by	486	No function assigned by TIGR
TIGR		487	putative phospholipase D-gamma
456	unknown protein	488	cell elongation protein, Dwarf1
457	unknown protein	489	germin-like protein
458	gamma-adaptin, putative	490	hevein-like protein precursor (PR-
459	UDP rhamnose		4)
	anthocyanidin-3-glucoside	491	rac-like GTP binding protein
	rhamnosyltransferase - like		(ARAC5)
	protein	492	phosphoprotein phosphatase, type
460	carbonate dehydratase - like		1 catalytic subunit
	protein	493	ubiquitin-protein ligase UBC9
461	putative microtubule-	494	xyloglucan endotransglycosylase-
	associated protein		related protein XTR-7
462	putative ribophorin I	495	cysteine synthase
463	putative zinc finger protein	496	putative villin 2
464	chloroplast FtsH protease,	497	glutathione S-transferase
	putative	498	5-adenylylsulfate reductase
465	putative protein	499	arginine decarboxylase
466	unknown protein	500	ATHP2, putative
467	putative LEA protein	501	ornithine carbamoyltransferase
468	putative protein	precui	
469	putative protein	502	puative protein
470	unknown protein	503	putative protein
471	putative purple acid	504	unknown protein
	phosphatase	505	putative protein
472	unknown protein	506	putative protein
473	putative protein	507	unknown protein
474	unknown protein	508	unknown protein
475	chlorophyll binding protein,	509	unknown protein
	putative	510	unknown protein
		511	hypothetical protein

512	putative protein	552	putative CCCH-type zinc finger
513	putative DnaJ protein		protein
514	plastocyanin	553	MAP kinase kinase 2
515	unknown protein	554	ethylene-insensitive3-like1 (EIL1)
516	unknown protein	555	histidine transport protein (PTR2-
517	unknown protein		B)
518	unknown protein	556	putative auxin-induced protein
519	unknown protein		AUX2-11
520	unknown protein	557	hydroxyacylglutathione hydrolase
521	putative ATP-dependent RNA helicase		cytoplasmic (glyoxalase II) (GLX II)
522	non-race specific disease	558	delta-8 sphingolipid desaturase
	resistance protein (NDR1)	559	cellulose synthase catalytic subunit
523	hypothetical protein		(Ath-A)
524	putative protein	560	nitrate transporter (NTL1)
525	putative protein	561	DNA-binding homeotic protein
526	putative protein		Athb-2
527	copper transport protein	562	hypothetical protein
528	putative protein	563	aspartate aminotransferase
529	unknown protein	564	4-coumarate:CoA ligase 1
530	unknown protein	565	pyruvate dehydrogenase E1 beta
531	unknown protein		subunit, putative
532	putative protein kinase	566	nucleotide diphosphate kinase Ia
533	unknown protein		(emb CAB58230.1)
534	putative protein	567	chloroplast Cpn21 protein
535	putative protein	568	ATP dependent copper transporter
536	hypothetical protein	569	very-long-chain fatty acid
537	putative protein		condensing enzyme (CUT1)
538	putative AP2 domain	570	putative purine-rich single-stranded
	transcription factor		DNA-binding protein
539	putative nitrilase	571	serine/threonine protein
540	putative protein		phosphatase (type 2A)
541	putative tetrahydrofolate	572	isopentenyl
	synthase		diphosphate:dimethylallyl
542	heat-shock protein		diphosphate isomerase (IPP2)
543	unkown protein	573	putative c2h2 zinc finger
544	unknown protein		transcription factor
545	histone H4	574	putative 20S proteasome beta
546	hypothetical protein		nit PBC2
547	unknown protein	575	nucleoside diphosphate kinase 3
548	putative protein	(ndpl	
549	predicted protein	576	ras-related small GTP-binding
550	putative dihydrolipoamide	prote	
	succinyltransferase	577	putative 4-coumarate:CoA ligase 2
551	actin 3		

578	transcription factor HBP-1b homolog (sp P43273)	609	photosystem II oxygen-evolving complex protein 3 - like
579	biotin synthase (Bio B)	610	sedoheptulose-bisphosphatase
580	homeobox protein HAT22	010	precursor
581	putative preprotein	611	glutathione S-transferase (GST6)
201	translocase SECY protein	612	geranylgeranyl reductase
582	carbamoylphosphate	613	hypothetical protein
362	synthetase, putative	614	hypothetical protein
583	putative protein kinase,	615	phosphoribulokinase precursor
ADK1	putative protein kinase,	616	high mobility group protein
584	putative nuclear DNA-	010	(HMG1), putative
384		617	protease inhibitor II
505	binding protein G2p	618	protease inhibitor II
585	hypothetical protein	619	cytochrome P450 90A1
586	hypothetical protein	619	(sp Q42569)
587	unknown protein	620	unknown protein
588	unknown protein	620	
589	molybdopterin synthase	621	heat shock protein 90 tubulin beta-9 chain
	(CNX2)		
590	putative ribosomal protein	623	putative ubiquitin carboxyl
L6			terminal hydrolase
591	unknown protein	624	protein kinase
592	En/Spm-like transposon	625	DRE/CRT-binding protein
proteir			DREB1C
593	putative protein	626	histidyl-tRNA synthetase
594	putative protein	627	splicing factor, putative
595	unknown protein	628	glutamyl-tRNA synthetase
596	hypothetical protein	629	putative RING zinc finger protein
597	unknown protein	630	phytochelatin synthase
598	unknown protein		(gb AAD41794.1)
599	putative lysosomal acid	631	putative C2H2-type zinc finger
lipase			protein
600	unknown protein	632	putative ligand-gated ion channel
601	unknown protein		protein
602	NifS-like aminotranfserase	633	putative ribosomal-protein S6
603	actin 8		kinase (ATPK6)
604	hypothetical protein	634	MOLYBDOPTERIN
605	putative protein		BIOSYNTHESIS CNX1
606	heat-shock protein (At-		PROTEIN
	hsc70-3)	635	temperature-sensitive omega-3
607	putative protein disulfide		fatty acid desaturase, chloroplast
	isomerase precursor		precursor (sp P48622)
608	adenosine nucleotide	636	adenylosuccinate synthetase
	translocator	637	putative 14-3-3 protein
		638	putative cytochrome P450

639	putative two-component	667	putative receptor-like protein
	response regulator 3 protein		kinase
640	putative RING-H2 zinc	668	putative disease resistance protein
	finger protein ATL6	669	receptor-like protein kinase - like
641	No function assigned by	670	ubiquitin activating enzyme 2
TIGR			(gb AAB37569.1)
642	small zinc finger-like	671	No function assigned by TIGR
protein		672	putative receptor-like protein
643	hypothetical protein		kinase
644	MAP kinase (ATMPK6)	673	K+ transporter, AKT1
645	vacuolar ATP synthase,	674	shaggy-like kinase beta
putativ	ve	675	heat shock protein 70
646	kinesin-like protein	676	plasma membrane intrinsic protein
647	serine/threonine-specific		1a
proteir	n kinase NAK	677	HSP90-like protein
648	No function assigned by	678	histone H1, putative
TIGR		679	unknown protein
649	ACTIN 2/7 (sp P53492)	680	dnaK-type molecular chaperone
650	phosphoglycerate kinase,		hsc70.1 - like
	putative	681	gamma-glutamylcysteine
651	homeotic protein BEL1		synthetase
	homolog	682	peroxidase (ATP22a)
652	proline iminopeptidase	683	putative serine carboxypeptidase
653	pasticcino 1		precursor
654	serine/threonine protein	684	putative dioxygenase
kinase	•	685	glucose transporter
655	cytochrome P450	686	NOI protein, nitrate-induced
	monooxygenase	687	putative protein
	(CYP71B4)	688	putative protein
656	No function assigned by	689	unknown protein
TIGR		690	putative photosystem I reaction
657	putative GDSL-motif		center subunit II precursor
	lipase/hydrolase	691	putative protein
658	putative protein	692	unknown protein
659	unknown protein	693	cobalamin biosynthesis protein
660	hypothetical protein	694	adenine nucleotide translocase
661	putative glycosylation	695	glutathione transferase, putative
enzyn	ne	696	putative 60S ribosomal protein L21
662	No function assigned by	697	cytochrome P450 like protein
TIGR		698	cytochrome b245 beta chain
663	No function assigned by		homolog RbohAp108, putative
TIGR		699	RNA helicase, DRH1
664	unknown protein	700	putative aldolase
665	putative ABC transporter	701	farnesyltransferase subunit A
666	nifU-like protein		(FTA)

702	No function assigned by	725	putative protein
TIGR		726	NBD-like protein
703	putative putative sister-		(gb AAD20643.1)
	chromatide cohesion	727	AtHVA22c
	protein	728	unknown protein
704	calcium-dependent protein	729	phytoene synthase
	kinase		(gb AAB65697.1)
705	serine/threonine protein	730	protein kinase (AME2/AFC1)
	phosphatase type 2A,	731	hypothetical protein
	putative	732	cyclin-dependent protein kinase-
706	40S ribosomal protein S28		like protein
	(sp P34789)	733	photosystem II stability/assembly
707	RNA polymerase subunit		factor HCF136 (sp O82660)
708	DNA-damage-	734	hypothetical protein
	repair/toleration protein	735	DNA binding-like protein
	DRT102	736	putative protein
709	putative C2H2-type zinc	737	chorismate mutase
	finger protein	738	putative LRR receptor protein
710	putative adenosine		kinase
,	phosphosulfate kinase	739	putative chalcone synthase
711	lipase	740	putative protein kinase
712	putative violaxanthin de-	741	replicase, putative
	epoxidase precursor	742	putative cysteine proteinase
	(U44133)	743	60S ribosomal protein L36
713	aromatic rich glycoprotein,	744	unknown protein
	putative	745	CLC-b chloride channel protein
714	putative fumarase	746	putative ribosomal protein S14
715	flavonol synthase (FLS)	747	histone H2B like protein
(splO9	96330)		(emb CAA69025.1)
716	response regulator 5,	748	60S ribosomal protein L2
putati	ve	749	60S ribosomal protein L15
717	sulfate transporter		homolog
718	putative floral homeotic	750	ribosomal protein S27
protei	n, AGL9	751	ribosomal protein
719	putative ethylene-inducible	752	60S ribosomal protein L12
	protein	753	60s ribosomal protein L34
720	C-8,7 sterol isomerase	754	putative ribosomal protein S10
721	TCH4 protein	755	drought-induced protein like
	(gb AAA92363.1)	756	blue copper-binding protein, 15K
722	hypothetical protein	,	(lamin)
723	putative urease accessory	757	calmodulin-like protein
	protein	758	putative protein
724	molybdopterin synthase	759	No function assigned by TIGR
	sulphurylase	760	alpha-mannosidase, putative
	(gb AAD18050.1)	761	uncoupling protein (ucp/PUMP)

		706	1
762	homeodomain - like protein	786	calcium-dependent protein kinase
763	ribosomal protein S18,		71196)
putati		787	phosphoinositide specific
764	similar to SOR1 from the		phospholipase C
	fungus Cercospora	788	similarity to S-domain receptor-
	nicotianae		like protein kinase, Zea mays
765	60S ribosomal protein L13,	789	mitosis-specific cyclin 1b
	BBC1 protein	790	4-coumarate:CoA ligase 3
766	50S ribosomal protein L24,	791	transcription factor IIB (TFIIB)
	chloroplast precursor	792	unknown protein
767	putative ribosomal protein	793	hypothetical protein
768	unknown protein	794	hypothetical protein
769	aspartate aminotransferase	795	sugar transporter like protein
	(AAT1)	796	putative trypsin inhibitor
770	potassium channel protein	797	unknown protein
	AtKC	798	putative multispanning membrane
771	unknown protein		protein
772	peroxisomal targeting	799	receptor-like kinase, putative
	signal type 2 receptor	800	putative inosine-5-monophosphate
773	putative protein		dehydrogenase
774	Ras-related GTP-binding	801	inosine-5'-monophosphate
	protein (ARA-4)		dehydrogenase, putative
775	S-receptor kinase homolog	802	amino acid permease 6
,,,	2 precursor		(emb CAA65051.1)
776	pathogenesis-related group	803	NADPH-ferrihemoprotein
,,,	5 protein, putative		reductase (ATR2)
777	Nitrilase 4 (splP46011)	804	putative WRKY-type DNA binding
778	biotin carboxyl carrier		protein
770	protein of acetyl-CoA	805	putative ankyrin
	carboxylase precursor	806	putative hexose transporter
	(BCCP) (sp Q42533)	807	aquaporin/MIP - like protein
779	photosystem I reaction	808	Ser/Thr protein kinase isolog
117	centre subunit psaN	809	pectate lyase like protein
	precursor (PSI-N)	810	putative 60S ribosomal protein L17
	(sp P49107)	811	putative protein
780	3(2),5-bisphosphate	812	unknown protein
780	nucleotidase	813	phenylalanine ammonia-lyase
781	high affinity Ca2+	814	putative cytochrome P450
antip		011	monooxygenase
782	putative cytoskeletal	815	ARR1 protein, putative
prote		816	putative bHLH transcription factor
783	putative peroxidase	817	aminomethyltransferase-like
784	respiratory burst oxidase	0.7	precursor protein
prote		818	purple acid phosphatase precursor
785	beta-glucosidase	0.0	r r r r r r
100	bem-gracostaase		

8	19	AP2 domain containing	844	mercaptopyruvate
		protein, putative		sulfurtransferase, putative
8	20	ubiquitin-conjugating	845	putative thiosulfate
		enzyme E2-21 kD 1		sulfurtransferase
		(ubiquitin-protein ligase 4)	846	dihydrolipoamide S-
		(ubiquitin carrier protein 4)		acetyltransferase
		(sp P42748)	847	auxin transport protein REH1,
0	21	translation initiation factor	0.,	putative
_	22	putative VAMP-associated	848	putative auxin transport protein
0	22	protein	849	apyrase (Atapy1)
0	23		850	root cap 1 (RCP1)
-		spermidine synthase,	851	hypothetical protein
	utativ		852	putative protein
	24	putative protein		predicted protein of unknown
	25	unknown protein	853	
	26	AtKAP alpha	functio	
8	27	glyceraldehyde-3-	854	hypothetical protein
		phosphate dehydrogenase,	855	hypothetical protein
		putative	856	hypothetical protein
8	328	putative poly(A) binding	857	putative aldehyde dehydrogenase
		protein	858	putative peroxidase
8	329	alpha-tubulin, putative	859	UDP-glucose 4-epimerase - like
8	330	serine/threonine-specific		protein
		protein kinase ATPK64	860	indole-3-acetate beta-
		(pir S20918)		glucosyltransferase like protein
8	331	putative aspartate-tRNA	861	putative beta-1,3-glucanase
1	igase		862	disease resistance protein-like
	332	ras-related small GTP-	863	putative respiratory burst oxidase
		binding protein RAB1c		protein B
5	333	cycloartenol synthase	864	ubiquitin-conjugating enzyme
	334	No function assigned by		UBC3
	ΓIGR		865	cytoplasmic aconitate hydratase
	335	cytochrome P450	866	NADPH oxidoreductase, putative
	836	GTPase AtRAB8	867	PROTEIN TRANSPORT
	837	3-phosphoserine		PROTEIN SEC61 GAMMA
		hatase		SUBUNIT -like
	910sp 838	transcription factor CRC	868	putative protein
	839	nuclear cap-binding	869	unknown protein
•	037	protein; CBP20	870	60S acidic ribosomal protein P2
		(gb AAD29697.1)	871	No function assigned by TIGR
	840	chloroplast membrane	872	1,4-alpha-glucan branching
•	840		072	enzyme protein soform SBE2.2
	0.41	protein (ALBINO3)		precursor
	841	biotin holocarboxylase	873	calcium binding protein (CaBP-22)
	0.40	synthetase	873 874	putative phosphoglucomutase
	842	expansin AtEx6	0/4	putative phosphogracomutase
	843	unknown protein		

875	shaggy-like protein kinase	901	putative RAS superfamily GTP-
	etha (EC 2.7.1)		binding protein
876	pyruvate decarboxylase	902	disease resistance protein-like
	(gb AAB16855.1)	903	protein kinase like protein
877	hypothetical protein	904	glucuronosyl transferase-like
878	putative protein kinase		protein
879	putative protein kinase	905	putative homeodomain
880	putative leucine		transcription factor
	aminopeptidase	906	putative flavonol reductase
881	probable cytochrome P450	907	putative protein
882	protein kinase 6-like protein	908	salt-tolerance protein
883	arginine methyltransferase	909	40S ribosomal protein S30
	(pam1)	910	putative bZIP transcription factor
884	MYB96 transcription	911	putative protein
	factor-like protein	912	putative cinnamoyl CoA reductase
885	putative protein	913	unknown protein
886	metal ion transporter	914	putative RNA-binding protein
887	No function assigned by	915	phosphatidylinositol synthase
TIGR		(PIS1)	
888	flax rust resistance protein,	916	unknown protein
	putative	917	hydroxyproline-rich glycoprotein
889	fructose-2,6-	homol	og
	bisphosphatase, putative	918	50S ribosomal protein L15,
890	exonuclease RRP41	chlore	plast precursor
891	squamosa promoter binding	919	unknown protein
	protein-like 2	920	putative YME1 ATP-dependant
	(emb CAB56576.1)		protease
892	putative squamosa-	921	unknown protein
	promoter binding protein	922	putative ribosomal protein L28
893	O-acetylserine(thiol) lyase,	923	unknown protein
	putative	924	putative protein
894	snoRNA	925	protein ch-42 precursor,
895	snoRNA		chloroplast
896	ferredoxin-NADP+	926	protein serine/threonine kinase,
reduct	tase		putative
897	H+-transporting ATP	927	beta-VPE
	synthase chain 9 - like	928	putative vacuolar sorting receptor
	protein	929	putative translation initiation factor
898	photosystem I subunit III		IF-2
	precursor, putative	930	predicted protein of unknown
899	photosystem I subunit VI		function
	precursor	931	putative protein
900	auxin-binding protein 1	932	hypothetical protein
	precursor	933	hypothetical protein
		934	phosphate transporter, putative

935	No function assigned by	961	unknown protein
TIGR	No function assigned by	962	unknown protein
936	1 -41	963	unknown protein
930	beta subunit of protein farnesyl transferase ERA1	964	myrosinase-associated protein,
027		964	
937	putative glutamate	0.65	putative
	decarboxylase	965	hypothetical protein
938	putative indole-3-acetate	966	hypothetical protein
	beta-glucosyltransferase	967	No function assigned by TIGR
939	putative receptor-like	968	unknown protein
	protein kinase	969	hypothetical protein
940	UDP-galactose 4-	970	LAX1 / AUX1 -like permease
	epimerase-like protein	971	putative UDP-N-
941	putative proliferating cell		acetylglucosaminedolichyl-
	nuclear antigen, PCNA		phosphate N-
942	ubiquitin conjugating		acetylglucosaminephosphotransfer
	enzyme E2 (UBC13)		ase
943	cyclophilin (CYP2)	972	chorismate mutase CM2
944	cystatin	973	inner mitochondrial membrane
	CAA03929.1)	,,,	protein
945	putative alcohol	974	DEF (CLA1) protein
	rogenase	975	decoy
946	acidic ribosomal protein p1	976	citrate synthase
947	glutathione transferase	977	myosin
947	AtGST 10	978	40S ribosomal protein S19
	(emb CAA10457.1)	979	ripening-related protein - like
948	putative tropinone	980	putative signal peptidase I
		981	methionyl-tRNA synthetase
reduct		701	(AtcpMetRS)
949	ZIP4, a putative zinc	982	ribosomal protein precursor - like
0.50	transporter	982	50S ribosomal protein L21
950	unknown protein	983	chloroplast precursor (CL21)
951	putative protein	004	
952	putative protein	984	putative MYB family transcription
953	putative C2H2-type zinc	factor	
	finger protein	985	cyclophilin - like protein
954	putative RING zinc finger	986	hypothetical protein
	protein	987	naringenin 3-dioxygenase like
955	putative microtubule-	protei	
	associated protein	988	WD-repeat protein -like protein
956	unknown protein	989	putative serine carboxypeptidase II
957	putative protein	990	prenyltransferase, putative
958	putative protein	991	putative ligand-gated ion channel
phosp	hatase-2c		protein
959	V-ATPase subunit G (vag2	992	clathrin adaptor medium chain
	gene)		protein MU1B, putative
960	hypothetical protein	993	No function assigned by TIGR
	* *		

004		1005	
994	putative Tall-like non-	1025	putative tropinone reductase
005	LTR retroelement protein	1026	signal response protein (GAI)
995	putative 3-isopropylmalate	1027	putative steroid sulfotransferase
006	dehydrogenase	1028	hypothetical protein
996	3-isopropylmalate	1029	nucleic acid binding protein - like
	dehydratase, small subunit	1030	putative protein
997	unknown protein	1031	blue copper binding protein
998	unknown protein	1032	farnesylated protein (ATFP6)
999	unknown protein	1033	unknown protein
1000	hypothetical protein	1034	putative PCF2-like DNA binding
1001	putative protein		protein
1002	No function assigned by	1035	teosinte branched1 - like protein
TIGR		1036	putative protein
1003	putative beta-glucosidase	1037	unknown protein
1004	putative pectate lyase A11	1038	unknown protein
1005	putative beta-glucosidase	1039	2-oxoglutarate dehydrogenase, E1
1006	HD-Zip protein		component
1007	putative ubiquitin	1040	unknown protein
	conjugating enzyme	1041	unknown protein
1008	homeobox-leucine zipper	1042	CCAAT-binding transcription
	protein-like		factor subunit A(CBF-A)
1009	cytochrome P450 like	1043	hypothetical protein
protei		1044	putative growth regulator protein
1010	putative cysteine proteinase	1045	putative presenilin
	inhibitor B (cystatin B)	1046	putative expansin
1011	ethylene response sensor	1047	ribosomal - like protein
(ERS)		1048	unknown protein
1012	putative SWH1 protein	1049	unknown protein
1013	putative glutathione S-	1050	putative protein
	transferase	1051	putative protein
1014	putative protein	1052	unknown protein
1015	unknown protein	1053	unknown protein
1016	putative protein	1054	unknown protein
	phosphatase 2C	1055	unknown protein
1017	dnaJ protein homolog atj3	1056	unknown protein
1018	ferredoxin	1057	putative protein
1019	hypothetical protein	1058	putative protein
1020	putative sugar transport	1059	argininosuccinate lyase (AtArgH)
	protein, ERD6	1060	disease resistance protein homolog
1021	putative DnaJ protein	1061	aldehyde dehydrogenase like
1022	putative AP2 domain	protei	
	transcription factor	1062	GBF2, G-box binding factor
1023	putative protein	1063	CDPK-related kinase
1024	putative cyclin-dependent	1064	endo-1,4-beta-glucanase
	kinase regulatory subunit	1065	putative serine protease

1066	serine/threonine-specific	1091	putative ATP-dependent RNA
	lecRK1 precursor,lectin		helicase
recepto		1092	putative protein
1067	putative MAP kinase	1093	putative HMG protein
1068	RNase L inhibitor-like	1094	squalene monooxygenase 2
protein			(squalene epoxidase 2) (SE 2)
1069	No function assigned by		(sp O65403)
TIGR		1095	eukaryotic peptide chain release
1070	AP2 domain transcription		factor subunit 1, putative
	factor	1096	auxin-induced protein - like
1071	polygalacturonase	1097	putative lipoamide dehydrogenase
	isoenzyme 1 beta subunit,	1098	putative protein
	putative	1099	unknown protein
1072	putative lipid transfer	1100	putative oligopeptide transporter
proteir	1	1101	putative translation elongation
1073	putative protein kinase		factor ts
1074	putative protein	1102	putative CCAAT-binding
1075	ATP-dependent RNA		transcription factor subunit
	helicase like protein	1103	putative ABC transporter
1076	putative cyclic nucleotide-	1104	putative superoxide-generating
	regulated ion channel		NADPH oxidase flavocytochrome
	protein	1105	aspartate kinase-homoserine
1077	COP1 like protein		dehydrogenase - like protein
1078	putative peroxidase	1106	putative bHLH transcription factor
1079	putative NAK-like ser/thr	1107	putative geranylgeranyl transferase
	protein kinase		type I beta subunit
1080	putative cytochrome C	1108	putative ARP2/3 protein complex
1081	cytochrome c		subunit p41
1082	putative serine	1109	sulphite reductase
	carboxypeptidase II	1110	putative auxin-regulated protein
1083	acyl-(acyl carrier protein)	1111	transcription factor scarecrow-like
	thioesterase		14, putative
1084	DNA-binding factor,	1112	unknown protein
putati	ve	1113	monooxygenase 2 (MO2)
1085	MAP3K delta-1 protein	1114	putative amine oxidase
kinase	•	1115	zinc finger protein, putative
1086	AtMlo-h1-like protein	1116	DNA-binding protein, putative
1087	No function assigned by	1117	putative protein
TIGR		1118	putative protein
1088	putative expansin	1119	Avr9 elicitor response like protein
1089	defender against cell death	1120	putative protein
	protein, putative	1121	hypothetical protein
1090	glycolate oxidase - like	1122	putative nucleotide-sugar
protei	n		dehydratase
-		1123	UFD1 like protein

1124	putative trans-	1155	cytochrome c oxidoreductase like
	transferase		protein
1125	outward rectifying	1156	putative
	potassium channel KCO		carboxymethylenebutenolidase
1126	unknown protein	1157	unknown protein
1127	putative	1158	unknown protein
	acetylesterase	1159	unknown protein
1128	putative protein	1160	unknown protein
1129	No function assigned by	1161	unknown protein
TIGR		1162	unknown protein
1130	unknown protein	1163	auxin-induced protein (IAA20)
1131	unknown protein	1164	50S ribosomal protein L4
1132	unknown protein	1165	putative DNA topoisomerase III
1133	protein phosphatase		beta
homol	og (PPH1)	1166	No function assigned by TIGR
1134	unknown protein	1167	isp4 like protein
1135	No function assigned by	1168	putative protein kinase
TIGR		1169	hypothetical protein
1136	unknown protein	1170	putative pyrophosphatefructose-
1137	unknown protein		6-phosphate 1-phosphotransferase
1138	unknown protein	1171	putative protein
1139	putative protein	1172	putative protein
1140	unknown protein	1173	putative protein
1141	putative ubiquinol	1174	unknown protein
	cytochrome-c reductase	1175	unknown protein
1142	unknown protein	1176	putative protein
1143	contains similarity to high-	1177	putative protein
	glucose-regulated protein 8	1178	unknown protein
	GB:AAF08813 GI:6449083	1179	unknown protein
	from [Homo sapiens]	1180	putative protein
1144	unknown protein	1181	brassinosteroid insensitive 1 gene
1145	putative cis-Golgi SNARE		(BRI1)
	protein	1182	putative receptor protein kinase
1146	unknown protein	1183	vacuolar-type H+-translocating
1147	glutamate-1-semialdehyde		inorganic pyrophosphatase
	aminotransferase	1184	protein kinase - like protein
1148	No function assigned by	1185	glycyl tRNA synthetase, putative
TIGR		1186	subtilisin proteinase - like
1149	hypothetical protein	1187	hypothetical protein
1150	unknown protein	1188	cytochrome P450-like protein
1151	unknown protein	1189	cytochrome p450 like protein
1152	unknown protein	1190	putative protein kinase
1153	scarecrow-like 3	1191	pectinesterase - like protein
1154	putative proline-rich protein	1192	putative receptor-like protein
			kinase

	1210	LE 1 (cor	it)
1193	peroxidase ATP17a -like	1219	putative AP2 domain transcription
1101	protein		factor
1194	No function assigned by	1220	brassinosteroid receptor kinase,
TIGR			putative
1195	cellulose synthase catalytic	1221	TINY-like protein
	subunit - like protein	1222	glucose-6-phosphate isomerase
1196	RAS-related protein, RAB7	1223	putative protein
1197	putative aspartate	1224	putative NAM (no apical
	aminotransferase		meristem)-like protein
1198	cyclophilin	1225	unknown protein
1199	putative SF2/ASF splicing	1226	putative nucleotide-binding protein
	modulator, Srp30	1227	bZIP transcription factor (POSF21)
1200	putative cytochrome b5	1228	ubiquitin activating enzyme - like
1201	glutamyl-tRNA reductase,		protein
	putative	1229	telomere repeat-binding protein
1202	putative MADS-box protein	1230	unknown protein
1203	ammonium transport	1231	mevalonate kinase
	protein (AMT1)	1232	putative protein
1204	No function assigned by	1233	hypothetical protein
TIGR		1234	disease resistance RPP5 like
1205	putative beta-ketoacyl-CoA		protein
syntha		1235	putative protein
1206	thaumatin-like protein	1236	putative pectinesterase
1207	putative methionine	1237	Ttg1 protein (emb CAB45372.1)
	peptidase	1238	FUSCA PROTEIN FUS6
1208	putative protein	1239	NHE1 Na+/H+ exchanger
	hatase 2C	1240	No function assigned by TIGR
1209	kinase-like protein	1241	Phospholipase like protein
1210	receptor-associated kinase	1242	unknown protein
isolog	-	1242	unknown protein
1211	mitochondrial ribosomal	1244	unknown protein
protei		1245	AUX1-like amino acid permease
	oleosin, 18.5K	1246	unknown protein
1212	chalcone isomerase	1247	putative C2H2-type zinc finger
1213	putative cyclin-dependent	1247	protein
1214	kinase regulatory subunit	1248	putative protein
1215	putative thaumatin-like	1248	putative protein
		1249	putative protein putative glucosyltransferase
protei		1251	putative glucosymansierase putative lipase
1216	putative two-component		
1017	response regulator protein	1252	putative protein
1217	TATA binding protein-	1253	putative thioredoxin
1010	associated factor, putative	1254	AIG2-like protein
1218	predicted protein of	1255	short-chain alcohol dehydrogenase
	unknown function	1055	like protein
		1256	hypothetical protein

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1257	putative protein	1287	No function assigned by TIGR
1258	putative protein	1288	serine/threonine protein kinase
1259	glutathione peroxidase -		ATPK10
	like protein	1289	putative lipase
1260	putative protein	1290	choline kinase GmCK2p -like
1261	putative disease resistance		protein
	response protein	1291	putative sugar transport protein,
1262	putative protein		ERD6
1263	senescence-associated	1292	MYB27 protein - like
	protein (SAG29)	1293	DNA-binding protein, putative
1264	glycolate oxidase, putative	1294	similar to cold acclimation protein
1265	extensin - like protein		WCOR413 [Triticum aestivum]
1266	putative protein	1295	unknown protein
1267	unknown protein	1296	aquaporin (plasma membrane
1268	putative disease resistance		intrinsic protein 2B)
	protein	1297	No function assigned by TIGR
1269	putative receptor-like	1298	P-Protein - like protein
	protein kinase	1299	No function assigned by TIGR
1270	putative receptor-like	1300	putative cytochrome P450
	protein kinase		monooxygenase
1271	basic chitinase	1301	putative cytochrome P450
1272	putative pectin		monooxygenase
	lesterase	1302	putative thioredoxin
1273	peroxidase ATP N	1303	stromal ascorbate peroxidase
1274	class 2 non-symbiotic	1304	ethylene responsive element
	hemoglobin		binding factor-like protein
1275	nitrate transporter		(AtERF6)
1276	Ca2+/H+-exchanging	1305	auxin transport protein EIR1
	protein-like		(gb AAC39513.1)
1277	putative protein	1306	putative CONSTANS-like B-box
1278	hydroxynitrile lyase like		zinc finger protein
proteir		1307	putative protein kinase
1279	putative AP2 domain	1308	mitochondrial Lon protease
	ription factor		homolog 1 precursor (sp O64948)
1280	pectin methylesterase,	1309	putative protein
putativ		1310	heme activated protein, putative
1281	putative protein	1311	putative cytochrome P450
1282	beta-glucosidase-like	1312	No function assigned by TIGR
proteii		1313	putative lipase
1283	CCAAT box binding factor/	1314	putative protein
	ription factor Hap2a	1315	putative sugar transporter protein
1284	putative fibrillin	1316	putative sucrose transport protein,
1285	xyloglucan endo-		SUC2
1006	transglycosylase	1317	putative protein
1286	putative 10kd chaperonin	1318	putative protein

1319	putative endochitinase	1351	unknown protein
1320	putative acetone-	1352	bZIP transcription factor - like
	cyanohydrin lyase	proteir	1
1321	putative protein	1353	Medicago nodulin N21-like protein
1322	calmodulin-like protein	1354	putative endo-1,4-beta glucanase
1323	hypothetical protein	1355	1-aminocyclopropane-1-
1324	cysteine proteinase like		carboxylate oxidase
proteir	1	1356	putative anion exchange protein
1325	heat shock protein 17.6-II	1357	SRG1-like protein
1326	heat shock protein 18	1358	putative protein
1327	Arabidopsis mitochondrion-	1359	putative phi-1-like phosphate-
	localized small heat shock		induced protein
	protein (AtHSP23.6-mito)	1360	putative protein
1328	unknown protein	1361	putative embryo-abundant protein
1329	putative WRKY-type DNA	1362	putative hydrolase
	binding protein	1363	unknown protein
1330	No function assigned by	1364	unknown protein
TIGR		1365	hexose transporter - like protein
1331	hypothetical protein	1366	unknown protein
1332	putative integral membrane	1367	unknown protein
	protein nodulin	1368	peptide transport - like protein
1333	putative protein	1369	unknown protein
1334	unknown protein	1370	putative peptide transporter
1335	3-isopropylmalate	1371	disease resistance protein, putative
	dehydratase, small subunit	1372	cysteine protease component of
1336	unknown protein		protease-inhibitor complex
1337	putative homeodomain	1373	putative cytochrome P450
	transcription factor	1374	putative protein
1338	unknown protein	1375	hypothetical protein
1339	putative protein	1376	unknown protein
1340	peroxidase ATP19a	1377	putative
1341	putative Na+/H+-		phosphoribosylaminoimidazolecar
	exchanging protein		boxamide formyltransferase
1342	putative auxin-regulated	1378	putative protein
	protein	1379	HSP like protein
1343	unknown protein	1380	unknown protein
1344	unknown protein	1381	unknown protein
1345	putative trehalose-6-	1382	putative cytochrome P450
	phosphate synthase	1383	similar to pectinesterase
1346	putative lectin	1384	putative glucosyltransferase
1347	Mlo protein-like	1385	thaumatin-like protein
1348	unknown protein	1386	drought-inducible cysteine
1349	ethylene response factor,		proteinase RD19A precursor
putati		1387	vegetative storage protein Vsp2
1350	unknown protein	1388	unknown protein

			/
1389	unknown protein	1417	G-box binding bZIP transcription
1390	anthranilate N-		factor
	benzoyltransferase - like	1418	putative protein
	protein	1419	putative protein
1391	delta-1-pyrroline 5-	1420	putative protein
	carboxylase synthetase	1421	ATFP4-like
	(P5C1)	1422	unknown protein
1392	glutathione S-conjugate	1423	unknown protein
	transporting ATPase	1424	putative protein
	(AtMRP1)	1425	invertase inhibitor homolog
1393	hypothetical protein	(emb C	CAA73335.1)
1394	hypothetical protein	1426	unknown protein
1395	unknown protein	1427	unknown protein
1396	putative protein	1428	putative cytochrome b5
1397	putative protein	1429	putative protein
1398	No function assigned by	1430	putative protein
TIGR		1431	putative protein
1399	unknown protein	1432	No function assigned by TIGR
1400	putative protein kinase	1433	putative copper/zinc superoxide
1401	unknown protein		dismutase
1402	hypothetical protein	1434	protein phosphatase ABI1
1403	unknown protein	1435	glutamate dehydrogenase 2
1404	putative calcium-binding	1436	No function assigned by TIGR
	EF-hand protein	1437	low-temperature-induced protein
1405	cinnamyl-alcohol		78 (sp Q06738)
	dehydrogenase ELI3-1	1438	putative myo-inositol 1-phosphate
1406	putative protein		synthase
1407	unknown protein	1439	phosphate transporter
1408	senescence-associated		(gb AAB17265.1)
	protein sen1	1440	4-hydroxyphenylpyruvate
1409	hypothetical protein		dioxygenase (HPD)
1410	putative cytochrome P450	1441	histone H1
1411	proline oxidase,	1442	hypothetical protein
	mitochondrial precursor	1443	No function assigned by TIGR
	(osmotic stress-induced	1444	neoxanthin cleavage enzyme-like
1.410	proline dehydrogenase)	1 4 4 5	protein
1412 3	putative response regulator	1445 1446	dehydration-induced protein RD22 zinc finger protein ZAT7
3 1413	hypothetical protein	1446	unknown protein
1414	glutamine-dependent	1448	unknown protein
1414	asparagine synthetase	1449	unknown protein
1415	lysine-ketoglutarate	1450	unknown protein
1713	reductase/saccharopine	1451	putative protein
1416	En/Spm-like transposon	1452	putative protein
protei		1453	RNA helicase, putative
protei	••	1 100	12.11110use, pumure

		(7
1454	putative glycine-rich	1483	unknown protein
proteir		1484	cold and ABA inducible protein
1455	hypothetical protein		kin1
1456	putative protein	1485	gamma-VPE (vacuolar processing
1457	peroxidase		enzyme)
1458	peroxidase ATP3a	1486	putative protein 1 photosystem II
	(emb CAA67340.1)		oxygen-evolving complex
1459	metallothionein-like protein	1487	myrosinase-associated protein,
1460	endomembrane-associated		putative
	protein	1488	transcription factor ATMYB4
1461	ferritin 1 precursor	1489	H-protein promoter binding factor-
1462	dehydrin RAB18-like		2a
	protein (sp P30185)	1490	ammonium transporter, puitative
1463	HSR201 like protein	1491	putative zeta-carotene desaturase
1464	light regulated protein,		precursor
putativ	re .	1492	high-affinity nitrate transporter
1465	Dr4(protease inhibitor)		NRT2
1466	mitogen activated protein	1493	light induced protein like
	kinase kinase (nMAPKK)	1494	putative AT-hook DNA-binding
1467	glutathione S-transferase	protein	ı
1468	transcriptional activator	1495	putative glycogenin
	CBF1/ CRT/CRE binding	1496	putative light repressible receptor
	factor 1		kinase
1469	homeobox-leucine zipper	1497	serine/threonine kinase - like
	protein ATHB-12	proteir	
1470	amino acid permease I	1498	putative peroxidase
1471	MAP kinase (ATMPK7)	1499	cytochrome P450 monooxygenase
1472	potassium channel protein	(CYP8	
	AKT3	1500	MYB-related transcription factor
1473	cytochrome P450		(CCA1)
	monooxygenase	1501	Terminal flower1 (TFL1)
	(CYP91A2)	1502	sulfate transporter ATST1
1474	putative transport protein	1503	RING-H2 finger protein RHA3b
1475	putative protein	1504	lipoxygenase, putative
1476	hypothetical protein	1505	serine O-acetyltransferase (EC
1477	putative protein		2.3.1.30) Sat-52 (pir S71207)
1478	hypothetical protein	1506	ferulate-5-hydroxylase (FAH1)
1479	receptor protein kinase-like	1507	En/Spm-like transposon protein,
1.400	protein	1500	putative
1480	serine/threonine protein	1508	calmodulin-binding - like protein
1.401	kinase - like protein	1509	hypothetical protein
1481	putative auxin-regulated	1510	somatic embryogenesis receptor-
1400	protein amino acid transport protein	1511	like kinase -like protein putative giberellin beta-
1482	AAP2	1311	hydroxylase
	AAF2		nyuroxyrase

		•	,
1512	putative pectinesterase	1542	60S acidic ribosomal protein P0
1513	putative protein	1543	putative protein
1514	unknown protein	1544	auxin-induced protein, putative
1515	ribosomal protein	1545	unknown protein
1516	low-temperature-induced	1546	hypothetical protein
	65 kD protein (sp Q04980)	1547	protein phosphatase 2C ABI2
1517	putative glucosyltransferase		(PP2C) (sp O04719)
1518	peroxidase	1548	peroxidase, prxr2
(emb C	AA67551.1)	1549	putative peroxidase ATP12a
1519	ankyrin-like protein	1550	putative beta-amylase
1520	ribosomal protein S11 - like	1551	putative acetone-cyanohydrin lyase
1521	hypothetical protein	1552	fatty acid elongase 3-ketoacyl-CoA
1522	glycoprotein(EP1), putative		synthase 1
1523	calnexin - like protein	1553	putative citrate synthase
1524	SRG1-like protein	1554	pEARLI 1-like protein
1525	ethylene response factor 1	1555	putative MYB family transcription
	(ERF1)		factor
1526	transcriptional activator	1556	putative transcription factor
	CBF1-like protein		MYB28
1527	xyloglucan endo-1,4-beta-	1557	RNA helicase-like protein
	D-glucanase (XTR-6)	1558	snoRNA
1528	putative cinnamyl alcohol	1559	putative protein kinase
	dehydrogenase	1560	growth regulator like protein
1529	gibberellin 3 beta-	1561	putative potassium transporter
	hydroxylase, putative	1562	putative protein
1530	auxin response transcription	1563	60S ribosomal protein L14
	factor 3 (ETTIN/ARF3)	1564	unknown protein
1531	No function assigned by	1565	putative RING-H2 zinc finger
TIGR		proteir	
1532	putative protein	1566	putative pollen surface protein
1533	similar to avrRpt2-induced	1567	unknown protein
	protein 1	1568	unknown protein
1534	unknown protein	1569	unknown protein
1535	hypothetical protein	1570	putative Ca2+-ATPase
1536	putative protein kinase	1571	1-aminocyclopropane-1-
1537	respiratory burst oxidase -		ylate synthase -like protein
	like protein	1572	putative beta-glucosidase
1538	glucose-6-	1573	transcription factor ZAP1
	phosphate/phosphate-	1574	oligopeptide transporter, putative
	translocator precursor,	1575	putative protein
	putative	1576	putative glucosyltransferase
1539	class 1 non-symbiotic	1577	putative serine/threonine kinase
	hemoglobin (AHB1)	1578	squalene epoxidase - like protein
1540	endochitinase isolog	1579	similar to 14KD proline-rich
1541	putative cytochrome P450		protein DC2.15 precursor

			,
	(sp P14009); similar to	1612	DnaJ-like protein
	ESTs emb Z17709 and	1613	putative inositol polyphosphate-5-
	emb Z47685		phosphatase
1580	unknown protein	1614	putative cytochrome P450
1581	unknown protein	1615	putative protein
1582	hypothetical protein	1616	unknown protein
1583	60S ribosomal protein L38	1617	putative protein
1584	flavin-containing	1618	hypothetical protein
	monooxygenase, putative	1619	putative protein
1585	remorin	1620	sucrose-UDP glucosyltransferase
1586	unknown protein	1621	glucose-6-phosphate 1-
1587	putative protein		dehydrogenase
1588	lipoxygenase	1622	unknown protein
1589	cold-regulated protein	1623	mitochondrial chaperonin (HSP60)
	COR6.6 (KIN2)	1624	sucrose transport protein SUC1
1590	Myb transcription factor	1625	putative protein disulfide isomerase
	homolog (ATR1)	1626	putative pollen-specific protein
1591	putative protein	1627	integral membrane protein,
1592	unknown protein		putative
1593	unknown protein	1628	rubredoxin, putative
1594	Ca2+-transporting ATPase	1629	putative protein
	- like protein	1630	disease resistance protein RPS4,
1595	protein phosphatase 2C		putative
	(AtP2C-HA)	1631	putative peptide/amino acid
1596	peroxidase ATP24a		transporter
1597	branched-chain alpha keto-	1632	peroxidase, putative
	acid dehydrogenase,	1633	ethylene receptor, putative (ETR2)
	putative	1634	protein phosphatase 2C (PP2C)
1598	putative beta-ketoacyl-CoA	1635	putative glutathione S-transferase
	synthase	1636	homeodomain transcription factor
1599	putative protein	(ATH	
1600	putative beta-galactosidase	1637	putative nitrate transporter
1601	putative protein	1638	putative ribosomal protein L9,
1602	60S ribosomal protein L27	cytoso	
1603	putative annexin	1639	putative DNA-binding protein
1604	NAC domain protein,	1640	beta-1,3-glucanase-like protein
putati		1641	putative zinc transporter
1605	unknown protein	1642	transcription factor TINY
1606	late embryogenesis	1643	putative aspartate kinase-
1.007	abundant protein LEA like		serine dehydrogenase
1607	unknown protein	1644	ethylene reponse factor-like AP2
1608	putative protein	1645	n transcription factor
1609	dehydrin Xero2	1645	peptide transporter - like protein trehalose-6-phosphate synthase like
1610 1611	putative zinc finger protein unknown protein	1040	protein
1011	unknown protein		protein

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1647	putative ribonuclease	1676	pathogenesis-related protein 1
1648	hypothetical protein		precursor, 19.3K
1649	putative DNA-binding	1677	R2R3-MYB transcription factor
proteir	n	1678	hypothetical protein
1650	nodulin-like protein	1679	putative chitinase
1651	trehalose-6-phosphate	1680	Mlo protein, putative
	phosphatase - like protein	1681	putative WRKY-type DNA binding
1652	succinate dehydrogenase		protein
	flavoprotein alpha subunit	1682	putative acyl-CoA synthetase
	(emb CAA05025.1)	1683	putative pathogenesis-related
1653	unknown protein		protein
1654	stress related protein,	1684	putative chitinase
putativ	ve	1685	germin precursor oxalate oxidase
1655	putative chloroplast	1686	endoxyloglucan transferase,
	initiation factor 3		putative
1656	putative protein	1687	putative protein
1657	hypothetical protein	1688	putative cytochrome P450
1658	putative CCCH-type zinc	1689	similar to Mlo proteins from H.
	finger protein		vulgare
1659	similar to harpin-induced	1690	putative tropinone reductase
	protein hin1 from tobacco	1691	extensin-like protein
1660	unknown protein	1692	putative sarcosine oxidase
1661	unknown protein	1693	putative protein
1662	hypothetical protein	1694	hypothetical protein
1663	No function assigned by	1695	late embryogenesis-abundant
TIGR			protein, putative
1664	putative protein	1696	beta-carotene hydroxylase
1665	putative glutathione S-	1697	putative calcium binding protein
	transferase TSI-1	1698	unknown protein
1666	putative protein	1699	unknown protein
1667	putative PTR2 family	1700	predicted glycosyl transferase
	peptide transporter	1701	hypothetical protein
1668	receptor kinase-like protein	1702	hypothetical protein
1669	putative sugar transport	1703	hypothetical protein
	protein, ERD6	1704	putative protein
1670	putative protein	1705	unknown protein
1671	nodulin-like protein	1706	putative protein
1672	unknown protein	1707	putative protein
1673	putative receptor-like	1708	serine/threonine kinase - like
	protein kinase		protein
1674	glutathione-conjugate	1709	No function assigned by TIGR
	transporter AtMRP4	1710	putative pectinesterase
1675	ascorbate oxidase-like	1711	peroxidase like protein
protei	n	1712	No function assigned by TIGR

1713	phenylalanine ammonia		Coenzyme A 3-O-
lyase (PAL1)		methyltransferase
1714	peroxidase	1740	disease resistance protein EDS1
(emb 0	CAA68212.1)	1741	putative protein kinase
1715	putative AMP deaminase	1742	Gluthatione reductase, chloroplast
1716	putative MYB family		precursor
transci	ription factor	1743	putative heat shock protein
1717	DNA-directed RNA	1744	aspartate kinase
polym	erase II, third largest subunit	1745	putative major intrinsic (channel)
1718	nucleotide pyrophosphatase		protein
	-like protein	1746	matrix metalloproteinase, putative
1719	putative peroxidase	1747	putative GDSL-motif
1720	calcium sensor homolog		lipase/hydrolase
	(gb AAC26110.1)	1748	putative protein
1721	putative GDSL-motif	1749	DAG-like protein
	lipase/hydrolase	1750	serine/threonine kinase -like
1722	putative nonspecific lipid-		protein
	transfer protein	1751	formamidase - like protein
1723	acyl-carrier protein (ACP),	1752	CER2
	putative	1753	26S proteasome subunit 4
1724	putative glycine	1754	pectinesterase like protein
dehyd	rogenase	1755	putative disease resistance protein
1725	AIG1	1756	putative RNA methyltransferase
1726	ACC synthase (AtACS-6)	1757	unknown protein
1727	cyclin delta-3	1758	HOMEOBOX PROTEIN
1728	putative RING zinc finger		KNOTTED-1 LIKE 4 (KNAT4)
	protein	1759	glycine-rich RNA-binding protein
1729	aldose 1-epimerase - like		AtGRP2 - like
	protein	1760	putative acetylornithine
1730	putative phospholipase		transaminase
1731	phosphoenolpyruvate	1761	putative Sec24-like COPII protein
	carboxylase	1762	putative berberine bridge enzyme
1732	putative galactinol synthase	1763	putative GH3-like protein
1733	unknown protein	1764	putative ABC transporter
1734	putative protein	1765	putative reticuline oxidase-like
1735	1-aminocyclopropane-1-		protein
	carboxylate oxidase	1766	pectate lyase - like protein
1736	thioredoxin (clone GIF1)	1767	protein disulfide-isomerase-like
	(pir S58118)		protein
1737	trehalose-6-phosphate	1768	putative protein
	phosphatase	1769	putative membrane transporter
1738	beta-1,3-glucanase 2 (BG2)	1770	unknown protein
. =	(PR-2)	1771	unknown protein
1739	putative S-adenosyl-L-	1772	putative RING-H2 zinc finger
	methionine:trans-caffeoyl-		protein

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1773	unknown protein	1807	glycine-rich RNA binding protein
1774	unknown protein		7
1775	unknown protein	1808	dehydrin, putative
1776	MADS-box protein	1809	putative endoxyloglucan
(AGL2	20)		glycosyltransferase
1777		1810	glutamate decarboxylase 1 (GAD
	amidophosphoribosyltransf		1) (sp Q42521)
erase 2	precursor	1811	delta 9 desaturase
1778	putative dihydrodipicolinate	1812	UDP-glucose glucosyltransferase
syntha	se	1813	CARBONIC ANHYDRASE 2
1779	hypothetical protein	1814	response reactor 2 (ATRR2)
1780	ABA-responsive protein -	1815	S-adenosyl-methionine-sterol-C-
like			methyltransferase, putative
1781	putative protein	1816	putative DNA-binding protein
1782	hypothetical protein		(RAV2-like)
1783	DNA-binding protein-like	1817	gamma glutamyl hydrolase,
1784	No function assigned by		putative
TIGR		1818	protein phosphatase - like
1785	transcription factor,	1819	unknown protein
putativ	re	1820	unknown protein
1786	nitrate reductase, putative	1821	unknown protein
1787	putative protein	1822	copper transport protein - like
1788	putative protein		protein
1789	putative protein	1823	hypothetical protein
1790	putative protein	1824	unknown protein
1791	unknown protein	1825	putative peptide methionine
1792	unknown protein		sulfoxide reductase
1793	tryptophan synthase beta-	1826	putative obtusifoliol 14-alpha
	subunit (TSB2)		demethylase
1794	hypothetical protein	1827	glutamate dehydrogenase (EC
1795	putative protein		1.4.1) 1 (pir S71217)
1796	putative DNA-binding	1828	unknown protein
proteir		1829	xyloglucan endo-1,4-beta-D-
1797	putative 40S ribosomal		glucanase precursor
	protein S10	1830	unknown protein
1798	putative protein	1831	SNF1 related protein kinase
1799	putative cytochrome P450		(ATSRPK1)
1800	putative protein	1832	putative protein
1801	putative protein	1833	putative chloroplast nucleoid DNA
1802	putative glucosyltransferase		binding protein
1803	No function assigned by	1834	hypothetical protein
TIGR		1835	putative protein
1804	putative protein	1836	putative thiamin biosynthesis
1805	putative protein		protein
1806	unknown protein	1837	unknown protein

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1838	unknown protein	1869	putative tyrosine aminotransferase
1839	putative RNA helicase	1870	thionin
1840	putative SF21 protein	1871	No function assigned by TIGR
	{Helianthus annuus}	1872	APETALA2 protein
1841	unknown protein	1873	MADS-box protein (AGL3)
1842	NBS/LRR disease	1874	putative monooxygenase
	resistance protein, putative	1875	ZFP3 zinc finger protein
1843	hypothetical protein	1876	cell division protein FtsZ
1844	unknown protein		chloroplast homolog precursor
1845	No function assigned by		(sp Q42545)
TIGR	υ.	1877	calreticulin, putative
1846	glycine-rich protein	1878	phosphoserine aminotransferase
(AtGR		1879	12-oxophytodienoate-10,11-
1847	No function assigned by		reductase
TIGR	ę,	1880	putative bHLH transcription factor
1848	putative protein	1881	pectin methylesterase (PMEU1),
1849	putative glucosyltransferase		putative
1850	hypothetical protein	1882	DNA-binding protein
1851	hypothetical protein	1883	carnitine racemase like protein
1852	putative protein	1884	putative protein
1853	putative disease resistance	1885	endoxyloglucan transferase
proteir	n		(dbj BAA81669.1)
1854	thaumatin, putative	1886	RMA1 RING zinc finger protein
1855	putative proline-rich protein	1887	ammonium transporter
1856	sterol-C-methyltransferase	1888	apyrase (gb AAF00612.1)
1857	superoxidase dismutase	1889	potassium uptake transporter - like
1858	TINY-like protein		protein
1859	calcium-dependent protein	1890	putative ABC transporter
kinase	e, putative	1891	potassium transporter-like protein
1860	hypothetical protein	1892	integral membrane protein,
1861	putative protein kinase		putative
1862	DNA-directed RNA	1893	putative protein
	erase (mitochondrial)	1894	pyruvate decarboxylase-1 (Pdc1)
1863	putaive DNA-binding	1895	putative malate oxidoreductase
protei		1896	putative histone H2B
1864	late embryogenesis	1897	snoRNA
	abundant M17 protein	1898	symbiosis-related like protein
1865	putative protein	1899	unknown protein
1866	delta-1-pyrroline-5-	1900	unknown protein
	carboxylate synthetase	1901	hypothetical protein
1867	putative 60s ribosomal	1902	putative protein
	protein L10	1903	copper-binding protein-like
1868	cytochrome P450	1904	putative protein
CYP8	6A1	1905	unknown protein
		1906	putative glyoxalase II

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1907	No function assigned by	1936	serine/threonine protein kinase.
TIGR		putativ	ve
1908	hypothetical protein	1937	potassium transporter - like protein
1909	flavanone 3-hydroxylase	1938	lactate dehydrogenase (LDH1)
(FH3)		1939	hypothetical protein
1910	putative laccase	1940	unknown protein
1911	putative protein kinase	1941	putative thaumatin
1912	myb-related protein, 33.3K	1942	putative reticuline oxidase-like
	(pir S71284)		protein
1913	unknown protein	1943	uracil phosphoribosyltransferase,
1914	endo-xyloglucan transferase		putative
	- like protein	1944	transcription factor, putative
1915	TMV resistance protein N -	1945	unknown protein
like	1	1946	unknown protein
1916	putative xyloglucan	1947	GATA transcription factor 4
	endotransglycosylase	1948	unknown protein
1917	unknown protein	1949	unknown protein
1918	proline transporter 2	1950	senescence-associated protein -like
1919	resistance protein, putative	1951	putative pollen allergen
1920	actin, putative	1952	unknown protein
1921	putative related to microbial	1953	putative protein
	divalent cation tolerance	1954	glycine-rich protein
	proteins	1955	putative protein
1922	unknown protein	1956	3-methyladenine DNA glycosylase,
1923	putative glycosyl		putative
transfe	rase	1957	endoplasmic reticulum-type
1924	unknown protein		calcium-transporting ATPase 4
1925	putative protein	1958	putative pectinesterase
	phosphatase 2C	1959	cytochrome P450-like protein
1926	unknown protein	1960	RNA-binding protein (cp33)
1927	serpin, putative	1961	CONSTANS-like 1
1928	cinnamyl-alcohol	1962	putative small heat shock protein
	ogenase CAD1	1963	hypothetical protein
1929	putative protein import	1964	unknown protein
recepto	or	1965	cytochrome P450 - like protein
1930	unknown protein	1966	cysteine proteinase inhibitor like
1931	unknown protein		protein
1932	putative protein	1967	nicotianamine synthase
1933	putative CDP-		(dbj BAA74589.1)
	glycerolglycerol-3-	1968	copper amine oxidase like protein
phosph			(fragment2)
	atidyltransferase	1969	putative SCARECROW gene
1934	unknown protein		regulator
1935	putative LRR receptor-like	1970	unknown protein
protein	kınase	1971	unknown protein

	1972	putative alanine acetyl	2001	auxin response factor 1
		transferase	2002	pathogenesis-related protein 1
	1973	unknown protein	precur	sor, 18.9K
	1974	unknown protein	2003	hypothetical protein
	1975	unknown protein	2004	unknown protein
	1976	putative extensin	2005	zinc finger protein Zat12
	1977	putative protein kinase	2006	unknown protein
	1978	putative protein kinase	2007	unknown protein
	1979	NADPH-dependent	2008	cyclin, putative
		codeinone reductase,	2009	2-dehydro-3-
		putative		phosphoheptonate aldolase
	1980	peroxidase	2010	glutathione synthetase gsh2
	1981	putative cytochrome P450	2011	heat shock protein 17
	1982	No function assigned by	2012	putative Na+-dependent inorganic
	TIGR	The range and application	2012	phosphate cotransporter
	1983	putative zinc-finger protein	2013	No function assigned by TIGR
		(B-box zinc finger domain)	2014	unknown protein
	1984	putative tyrosine	2015	putative protein
		aminotransferase	2016	similar to RING-H2 finger protein
	1985	hypothetical protein		RHC1a GB:AAC69854
	1986	DNA binding protein		GI:3790583 from [Arabidopsis
	1987	putative fatty acid elongase		thaliana]
	1988	bZIP transcription factor -	2017	calcium-binding protein - like
		like protein	2018	putative protein
	1989	xyloglucan	2019	putative aldehyde dehydrogenase
		fucosyltransferase, putative	2020	auxin-responsive GH3 - like
	1990	unknown protein		protein
	1991	unknown protein	2021	putative protein
	1992	putative protein	2022	Phosphoglycerate dehydrogenase -
	1993	myb factor, putative		like protein
	1994	Myb-family transcription	2023	unknown protein
		factor, putative	2024	unknown protein
	1995	putative fructose	2025	PSI type III chlorophyll a/b-
		bisphosphate aldolase		binding protein, putative
	1996	myrosinase-associated	2026	putative protein
		protein, putative	2027	putative protein
	1997	cytochrome P450 like	2028	glutaredoxin, putative
	proteir	1	2029	hypothetical protein
	1998	similar to SOR1 from the	2030	No function assigned by TIGR
		fungus Cercospora	2031	putative protein
		nicotianae	2032	jasmonate inducible protein,
	1999	similar to embryo-abundant		putative
		GB:L47672 GI:1350530	2033	putative polygalacuronase
		Picea glauca]		isoenzyme 1 beta subunit
	2000	alcohol dehydrogenase	2034	putative small heat shock protein

	1,101	L X (CO)	
2035	unknown protein	2068	putative chlorophyll A-B binding
2036	putative disease resistance	2000	protein
	protein	2069	Lhcb3 chlorophyll a/b binding
2037	putative protein	2007	protein (gb AAD28773.1)
2038	ethylene-responsive	2070	luminal binding protein
	element binding factor,		AA13948.1)
	putative	2071	hydroxypyruvate reductase (HPR)
2039	putative protein	2072	epoxide hydrolase (ATsEH)
2040	Pollen-specific protein	2073	putative protein (fragment)
	precursor like	2074	unknown protein
2041	putative protein	2075	hypothetical protein
2042	unknown protein	2076	putative glucosyl transferase
2043	EF-Hand containing protein	2077	putative glucosyl transferase
	-like	2078	putative 3-methylcrotonyl-CoA
2044	unknown protein	carbo	
2045	puative calcium-	2079	putative peroxidase
	transporting ATPase	2080	acyl-CoA oxidase
2046	antifungal protein-like		AC13497.1)
	(PDF1.2)	2081	alternative oxidase 1a precursor
2047	pathogenesis-related PR-1-	2082	putative transcription factor
	like protein		(MYB4)
2048	similar to MIo proteins	2083	serine acetyltransferase
	from H. vulgare	2084	ATP-sulfurylase
2049	putative steroid	2085	calreticulin (crt1)
sulfoti	ransferase	2086	putative prohibitin 2
2050	trehalase - like protein	2087	putative monodehydroascorbate
2051	thioredoxin fl		reductase
2052	unknown protein	2088	branched-chain alpha-keto acid
2053	alanine-glyoxylate		decarboxylase E1 beta subunit
	aminotransferase	2089	cytokinin oxidase - like protein
2054	integral membrane protein,	2090	putative receptor-like protein
	putative		kinase
2055	hypothetical protein	2091	unknown protein
2056	unknown protein	2092	hypothetical protein
2057	hypothetical protein	2093	No function assigned by TIGR
2058	unknown protein	2094	putative APG protein
2059	unknown protein	2095	glutathione S-transferase, putative
2060	unknown protein	2096	phytochrome-associated protein 1
2061	drought-induced-19-like 1		(PAP1)
2062	unknown protein	2097	amidophosphoribosyltransferase
2063	putative protein	2098	nonphototropic hypocotyl 1
2064	putative protein	2099	3-keto-acyl-CoA thiolase 2
2065	AIG2-like protein		(gb AAC17877.1)
2066	Lhca2 protein	2100	pEARLI 1
2067	phytocyanin	2101	glutathione reductase, cytosolic

		ZZ Z (4011	-7
2102	putative protein	2128	putative protein disulfide-
2103	putative protein		isomerase
2104	putative aldehyde oxidase	2129	unknown protein
2105	probable photosystem I	2130	beta-1,3-glucanase class I
	chain XI precursor		precursor
2106	photosystem II polypeptide,	2131	homeobox-leucine zipper protein
	putative		HAT5 (HD-ZIP protein 5) (HD-
2107	photosystem II reaction		ZIP protein ATHB-1)
	center 6.1KD protein	2132	putative cyclic nucleotide-
2108	33 kDa polypeptide of		regulated ion channel protein
	oxygen-evolving complex	2133	P II nitrogen sensing protein GLB I
	(OEC) in photosystem II	2134	H-protein promoter binding factor-
	(emb CAA75629.1)		1 (gb AAC24592.1)
2109	60S ribosomal protein	2135	GAST1-like protein
L11B	•	2136	cytochrome P450 GA3
2110	extA (emb CAA47807.1)	2137	putative protein
2111	zinc finger protein OBP4 -	2138	Myb-related transcription factor-
like	5 1	like pro	
2112	sterol delta7 reductase	2139	putative phloem-specific lectin
2113	putative RAS-related	2140	protein kinase - like protein
	protein, RAB11C	2141	unknown protein
2114	glucosyltransferase like	2142	SCARECROW transcriptional
proteir	1	regulat	or-like
2115	zinc finger protein (PMZ),	2143	unknown protein
	putative	2144	unknown protein
2116	6,7-dimethyl-8-	2145	putative protein
	ribityllumazine synthase	2146	calnexin homolog
	precursor	2147	PP1/PP2A phosphatases
2117	putative protein		opic regulator PRL2
2118	osmotin precursor	2148	xyloglucan endotransglycosylase,
2119	No function assigned by	putativ	
TIGR		2149	putative calmodulin
2120	ferredoxin precusor isolog	2150	spermine synthase (ACL5)
2121	GH3 like protein	2151	snoRNA
2122	non-specific lipid transfer	2152	photosystem I subunit V precursor,
	protein		putative
2123	homeodomain transcription	2153	putative potassium transporter
	factor (HAT9)	2154	Homeodomain - like protein
2124	putative cytochrome P450	2155	putative protein
	monooxygenase	2156	unknown protein
2125	putative protein kinase	2157	CALMODULIN-RELATED
2126	putative protein		PROTEIN 2, TOUCH-INDUCED
2127	glyceraldehyde-3-		(TCH2)
	phosphate dehydrogenase	2158	putative protein phosphatase 2C

2159	monosaccharide transport	2187	defender against cell death protein
	protein, STP4	2188	AP2 domain containing protein,
2160	hypothetical protein		putative
2161	unknown protein	2189	actin depolymerizing factor - like
2162	hypothetical protein		protein
2163	putative protein kinase	2190	putative calcium-dependent protein
2164	putative serine/threonine		kinase (U90439)
	protein kinase	2191	phosphoribosylanthranilate
2165	jasmonate inducible		transferase, putative
	protein, putative	2192	oligopeptide transporter, putative
2166	similar to several small	2193	calmodulin-like protein
	proteins (~100 aa) that are	2194	putative protease inhibitor
	induced by heat, auxin,	2195	MAP kinase
	ethylene and wounding	2196	DNA binding protein MybSt1,
	such as Phaseolus aureus		putative
	indole-3-acetic acid	2197	putative protein
	induced protein ARG	2198	putative protein
	(SW:32292)	2199	unknown protein
2167	unknown protein	2200	unknown protein
2168	MYB-like protein	2201	unknown protein
2169	putative protein kinase	2202	putative protein
2170	unknown protein	2203	unknown protein
2171	CLC-d chloride channel	2204	unknown protein
proteir	1	2205	hypothetical protein
2172	cytochrome P450-like	2206	uncharacterized protein
proteir		2207	putative protein
2173	putative glutathione S-	2208	hypothetical protein
	transferase	2209	peroxidase (emb CAA66967.1)
2174	putative mandelonitrile	2210	putative flavonol 3-O-
lyase	F	glucos	yltransferase
2175	hypothetical protein	2211	putative flavonol 3-O-
2176	putative trypsin inhibitor	glucos	yltransferase
2177	male sterility 2-like protein	2212	putative protein
	(emb CAA68191.1)	2213	glycerol-3-phosphate
2178	unknown protein		ansferase
2179	unknown protein	2214	putative beta-1,3-glucanase
2180	putative protein	2215	putative ethylene response element
2181	putative peroxidase		g protein (EREBP)
2182	putative thromboxane-A	2216	putative CONSTANS-like B-box
2102	synthase		nger protein
2183	putative cytochrome P450	2217	putative protein
2184	peroxidase ATP21a	2218	unknown protein
2185	unknown protein	2219	putative trehalose-6-phosphate
2186	putative glutathione S-		hatase (AtTPPA)
2100	transferase	2220	putative protein

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2221	putative protein	2251	lysine and histidine specific
2222	unknown protein		transporter, putative
2223	unknown prptein	2252	putative protein
2224	unknown protein	2253	putative protein
2225	hypothetical protein	2254	putative sugar transporter protein
2226	putative metal-binding	2255	12S cruciferin seed storage protein
proteir		2256	putative auxin-induced protein,
2227	putative	2200	IAA17/AXR3-1
	phosphoribosylglycinamide	2257	putative cyclin D
	synthetase	2258	farnesyl diphosphate synthase
2228	unknown protein		precursor (gb AAB49290.1)
2229	putative protein	2259	putative potassium transport
2230	unknown protein	220,	protein (TRH1)
2231	unknown protein	2260	putative NPK1-related MAP kinase
2232	putative beta-galactosidase	2261	putative protein
2233	putative protein kinase	2262	putative ABC transporter
2234	putative protein	2263	putative DNA-directed RNA
2235	putative protein		polymerase subunit
2233	phosphatase 2C	2264	putative small nuclear
2236	putative growth regulator		ribonucleoprotein E
	protein	2265	unknown protein
2237	putative ABC transporter	2266	reticuline oxidase - like protein
2238	chloride channel	2267	putative 1-aminocyclopropane-1-
	(emb CAA70310.1)		carboxylate oxidase
2239	adrenodoxin - like protein	2268	similar to MIo proteins from H.
2240	NAM (no apical meristem)-		vulgare
	like protein	2269	long-chain-fatty-acidCoA ligase-
2241	putative transcription factor		like protein
	MYB41	2270	putative protein
2242	Myb DNA binding protein -	2271	chromatin remodelling complex
like			ATPase chain ISWI -like protein
2243	AtMYB84	2272	hypothetical protein
2244	photosystem II type I	2273	latex-abundant protein, putative
	chlorophyll a/b binding	2274	N-acetylornithine deacetylase-like
	protein		protein, fragment
2245	putative aspartic proteinase	2275	putative DNA-binding protein
2246	jasmonate inducible	2276	putative anthranilate N-
	protein, putative		hydroxycinnamoyl/benzoyltransfer
2247	putative protein	2277	ase
2248	No function assigned by	2277	putative DNA binding protein
TIGR		2278	cytochrome P450 - like protein
2249	putative phosphatidylserine	2279 2280	putative DNA-binding protein putative peptide transporter
2250	synthase	2280	putative peptide transporter putative reticuline oxidase-like
2250	putative nicotianamine	2281 protei	
	synthase	protei	11

22	82	thioredoxin, putative	2313	putative protein kinase
	2283 nodulin-like protein		2314	indoleacetic acid (IAA)-inducible
22		UDP-galactose transporter -		gene (IAA7)
		otein	2315	ATP-dependent Clp protease
22		putative fibrillin		regulatory subunit CLPX
22		unknown protein	2316	DNA-binding protein RAV1
22		unknown protein	2317	putative protein
22		unknown protein	2318	hypothetical protein
22		hypothetical protein	2319	unknown protein
22	90	glyceraldehyde 3-phosphate	2320	unknown protein
		dehydrogenase A subunit	2321	putative protein
		(GapA)	2322	putative thioredoxin reductase
22	91	predicted protein of	2323	unknown protein
		unknown function	2324	putative lectin
22		putative protein	2325	No function assigned by TIGR
	93	putative protein	2326	beta-fructosidase
22	94	myb-like protein	2327	chlorophyll a/b-binding protein
	95	hypothetical protein		CP29
22	96	putative U5 small nuclear	2328	photosystem I subunit PSI-E - like
		ribonucleoprotein, an RNA		protein
		helicase	2329	peroxidase ATP8a
22	97	unknown protein	2330	putative fructose bisphosphate
22	98	cinnamyl alcohol		aldolase
		dehydrogenase - like	2331	zinc finger protein ATZF1,
		protein		putative
22	99	hypothetical protein similar	2332	DegP protease precursor
		to extensin-like protein	2333	transcription factor-like protein
	00	unknown protein	2334	calcium-dependent protein kinase
23	01	putative chlorophyll a/b	2335	hypothetical protein
		binding protein	2336	putative protein
23	02	probable plasma membrane	2337	glucose-1-phosphate
		intrinsic protein 1c		adenylyltransferase (APL3)
23	03	hexokinase (ATHXK2)	2338	No function assigned by TIGR
23	04	calcium-dependent protein	2339	putative Eukaryotic initiation factor
		kinase		4A
23	05	5'-adenylylphosphosulfate	2340	No function assigned by TIGR
		reductase, putative	2341	unknown protein
23	06	Erd1 protein precursor	2342	beta tubulin 1, putative
		(sp P42762)	2343	one helix protein (OHP)
	07	putative protein	2344	No function assigned by TIGR
	808	putative protein	2345	zinc finger protein 5, ZFP5
	09	unknown protein	2346	putative MYB family transcription
23	310	BCS1 protein-like protein		factor
23	11	putative protein	2347	putative amino acid transporter
23	312	putative protein		protein

2348	putative potassium	2374	putative PHD-type zinc finger
transp 2349			protein
2350	protein kinase (AFC2)	2375	nuclear RNA binding protein A-
	putative protein		like protein
2351 TLCD	No function assigned by	2376	unknown protein
TIGR		2377	unknown protein
2352	putative ubiquitin-	2378	unknown protein
	gating enzyme E2	2379	putative amino-cyclopropane-
2353	unknown protein		carboxylic acid oxidase (ACC
2354	cytochrome P450		oxidase)
	oxygenase (CYP71B3)	2380	hypothetical protein
2355	putative myrosinase-	2381	indole-3-acetate beta-
	ng protein		glucosyltransferase like protein
2356	putative vacuolar sorting	2382	predicted protein
recept		2383	unknown protein
2357	uridine diphosphate glucose	2384	No function assigned by TIGR
epime	rase	2385	putative photosystem I reaction
2358	shaggy related protein		center subunit IV
	e, ASK-GAMMA	2386	putative homeodomain
2359	ankyrin repeat protein		transcription factor
EMB5	506	2387	putative purple acid phosphatase
2360	putative beta-alanine-		precursor
	pyruvate aminotransferase	2388	No function assigned by TIGR
2361	putative alcohol	2389	nitrate reductase 1 (NR1)
dehyd	rogenase	2390	putative casein kinase II beta
2362	putative receptor-like		subunit
	protein kinase	2391	pEARLI 1-like protein
2363	unknown protein	2392	putative protein
2364	putative methylmalonate	2393	No function assigned by TIGR
	semi-aldehyde	2394	unknown protein
	dehydrogenase	2395	putative cell wall-plasma
2365	hypothetical protein		membrane disconnecting CLCT
2366	unknown protein		protein (AIR1A)
2367	peroxidase ATP13a	2396	unknown protein
2368	putative glutathione	2397	scarecrow-like 11 - like
peroxi	dase	2398	putative anthocyanidin synthase
2369	squamosa promoter binding	2399	putative AP2 domain transcription
	protein-like 7		factor
2370	photosystem II core	2400	caffeoyl-CoA O-methyltransferase
	complex protein, putative		- like protein
2371	snoRNA	2401	unknown protein
2372	photosystem I subunit X	2402	putative protein kinase
	precursor	2403	cytochrome P450 -like protein
2373	MYB transcription factor	2404	putative MADS-box protein ANR1
	(Atmyb2)	2405	putative glutathione S-transferase
			. B

hypothetical protein	2437	putative protein
similar to gibberellin-	2438	unknown protein
regulated proteins	2439	unknown protein
unknown protein	2440	putative protein
putative sensory	2441	No function assigned by TIGR
transduction histidine	2442	MADS-box protein AGL14
kinase	2443	No function assigned by TIGR
similar to late	2444	peptidylprolyl isomerase
embryogenesis abundant	2445	putative s-adenosylmethionine
proteins		synthetase
unknown protein	2446	peroxidase
putative protein	2447	ferrochelatase-I
	2448	putative eukaryotic initiation factor
RNA helicase		4, eIF4
putative protein	2449	drought-inducible cysteine
putative sucrose synthetase		proteinase RD21A precursor -like
beta-fructofuranosidase 1		protein
putative indole-3-acetate	2450	unknown protein
lucosyltransferase	2451	unknown protein
hypothetical protein	2452	No function assigned by TIGR
DNA-directed RNA	2453	No function assigned by TIGR
erase II, third largest subunit	2454	salt-inducible like protein
	2455	glucose-6-phosphate 1-
		dehydrogenase
	2456	3-hydroxy-3-methylglutaryl CoA
		reductase (AA 1-592)
		hypothetical protein
		putative protein
	2459	putative putative 60S ribosomal
		protein L17
		putative inorganic pyrophosphatase
	2461	putative gamma-
		glutamyltransferase
	2462	heat shock transcription factor -
		like protein
		mitochondrial chaperonin hsp60
		unknown protein
	2465	putative zinc finger protein
		identical to T10M13.22
		putative uridylyl transferase
		nodulin-like protein
		putative B-box zinc finger protein
		No function assigned by TIGR
	2470	putative metalloproteinase
unknown protein		
	similar to gibberellin- regulated proteins unknown protein putative sensory transduction histidine kinase similar to late embryogenesis abundant proteins unknown protein putative Protein putative ATP-dependent RNA helicase putative protein putative sucrose synthetase beta-fructofuranosidase 1 putative indole-3-acetate lucosyltransferase hypothetical protein DNA-directed RNA	similar to gibberellin- regulated proteins 2438 regulated proteins 2439 unknown protein 2440 putative sensory 2441 transduction histidine 2442 kinase 2443 similar to late 2444 embryogenesis abundant 2445 proteins 2446 putative protein 2447 putative ATP-dependent 2448 RNA helicase putative protein 2449 putative sucrose synthetase beta-fructofuranosidase 1 putative indole-3-acetate dlucosyltransferase 2451 hypothetical protein 2452 DNA-directed RNA 2453 erase II, third largest subunit putative transcription factor homeobox-leucine zipper n ATHB-5 (HD-zip protein 2456 n ATHB-5 (HD-zip protein 3-5) (splP46667) putative ftsH chloroplast 2458 replication protein A1 - like hypothetical protein 2461 unknown protein 2461 putative methionine 2461 muknown protein 2461 putative delongase - like 2463 protein (cer2-like) 2465 unknown protein 2465 putative disease resistance response protein 2468 putative protein 2468 putative protein 2468 putative protein 2469 putative protein 2460 putative protein 2468 putative protein 2468 putative protein 2468 putative protein 2469 putative protein 2460 putative protein 2468 putative protein 2468 putative protein 2469 putative protein 2460

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2471	putative cellular apoptosis	2504	unknown protein
	susceptibility protein	2505	unknown protein
2472	hypothetical protein	2506	60S ribosomal protein L10A
2473	hypothetical protein	2507	putative protein
2474	scarecrow-like 13 (SCL13)	2508	receptor protein kinase (IRK1),
2475	putative nucleoside		putative
	triphosphatase	2509	putative nematode-resistance
2476	unknown protein		protein
2477	No function assigned by	2510	tubulin alpha-5 chain-like protein
TIGR		2511	putative DNA-binding protein
2478	hypothetical protein	2512	unknown protein
2479	putative phospholipase	2513	putative RGA1, giberellin repsonse
2480	putative snRNP protein		modulation protein
2481	putative protein	2514	non phototropic hypocotyl 1-like
2482	putative lipase	2515	RING-H2 finger protein RHA1b
2483	putative nonsense-mediated	2516	putative myb-protein
	mRNA decay protein	2517	hydroperoxide lyase (HPOL) like
2484	No function assigned by		protein
TIGR		2518	serine/threonine-protein kinase,
2485	protochlorophyllide		PK7
	reductase precursor	2519	putative vacuolar proton-ATPase
2486	No function assigned by		subunit
TIGR		2520	putative polygalacturonase
2487	trehalose-6-phosphate	2521	putative ribosomal protein L8
0.400	synthase, putative	2522	putative adenylate kinase
2488	unknown protein	2523	germin-like protein (GLP10)
2489	germin-like protein	2524	putative chlorophyll a/b binding
2490	plastid protein		protein
2491	putative protein	2525	chloroplast single subunit DNA-
2492 2493	hypothetical protein	0.000	dependent RNA polymerase
2493	unknown protein	2526	putative protein
2494	unknown protein	2527	hypothetical protein
protein	histone deacetylase-like	2528	hypothetical protein
2496	unknown protein	2529	b-keto acyl reductase, putative
2497	unknown protein	2530 2531	cellulose synthase catalytic subunit
2498	putative protein	2551	putative 1-aminocyclopropane-1-
2499	putative protein	2532	carboxylate oxidase
2500	No function assigned by	2533	S-linalool synthase, putative phosphoribosyl-ATP
TIGR	140 Idiletion assigned by	2333	pyrophosphohydrolase (At-IE)
2501	putative zinc transporter	2534	disease resistance RPP5 like
ZIP2 -		2334	protein (fragment)
2502	unknown protein	2535	putative protein
2503	putative ribosomal-protein	2536	beta-galactosidase like protein
	S6 kinase (ATPK19)	2330	sem garactosidase like protein
	(//////////////////////////		

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2537	putative translation	2566	unknown protein
	initiation factor eIF-2,	2567	unknown protein
	gamma subunit	2568	unknown protein
2538	ankyrin like protein	2569	serine/threonine kinase - like
2539	histone H2A- like protein	protein	
2540	putative protein	2570	peroxidase (emb CAA66960.1)
2541	salt-tolerance zinc finger	2571	putative protein
	protein	2572	hypothetical protein
2542	unknown protein	2573	glycine-rich protein 2 (GRP2)
2543	putative protein	2574	unknown protein
2544	fructose-bisphosphate	2575	berberine bridge enzyme-like
aldola		protei	
2545	peroxidase	2576	unknown protein
	CAA66964.1)	2577	putative WD-repeat protein
2546	patatin-like protein	2578	serine/threonine kinase - like
2547	salt-inducible protein	2376	
homo		2579	protein
2548	hypothetical protein	2319	serine /threonine kinase - like
2549	xyloglucan endo-	2580	protein
2349	transglycosylase-like	2380	Cu2+-transporting ATPase-like protein
	protein	2581	
2550	trihelix DNA-binding	2582	translation initiation factor eIF4E
2330	protein (GT2)	2583	O-methyltransferase - like protein
2551	ubiquitin-conjugating	2383	translation initiation factor eIF3 -
2331	enzyme 16, putative	2594	like protein
2552	homeobox protein	2584	No function assigned by TIGR
2553	envelope Ca2+-ATPase	2585	unknown protein
2554	snap25a	2586 2587	hypothetical protein
2555	putative annexin	2587	unknown protein
2556	putative annexin	2589	unknown protein
2557	homeodomain transcription		glycine-rich protein like
2331	factor (ATHB-14)	2590 2591	putative disease resistance protein
2558	heat shock protein, putative	2591	putative Na+/Ca2+ antiporter
2559	peroxidase ATP23a	2392	putative hydroxymethylglutaryl-
2560	p68 RNA helicase, putative	2593	CoA lyase
2561	potassium transporter,	2393	putative
putativ			phosphoribosylaminoimidazole
2562	putative eukaryotic	2594	carboxylase
	ation initiation factor 2 alpha		SAR DNA-binding protein - like
	it, eIF2	2595 2596	response regulator, putative
2563	hypothetical protein	2596	fibrillin precursor-like protein
2564	carnitine racemase like	2397	beta-ketoacyl-CoA synthase
protei		2598	(FIDDLEHEAD) lectin like protein
2565	No function assigned by	2598	No function assigned by TIGR
TIGR	110 function assigned by	2399	ino function assigned by TIGR
AUL			

2600	acidic endochitinase	2629	vel
2000	(dbi BAA21861.1)	2630	unknown protein unknown protein
2601	unknown protein	2631	
2602	hypothetical protein	2632	unknown protein
2603	predicted OR23 protein of		nucleosome assembly protein I-like
2003		protei	
2604	unknown function	2633	membrane channel like protein
	putative protein	2634	anthocyanin2, putative
2605	hypothetical protein	2635	TWIN SISTER OF FT (TSF)
2606	glycerol-3-phosphate	2636	putative myb-related transcription
2505	dehydrogenase	factor	
2607	hypothetical protein	2637	hypothetical protein
2608	tat-binding protein, putative	2638	putative RING zinc finger protein
2609	putative protein	2639	amino acid transport protein AAT1
2610	putative trehalose-6-	2640	putative protein
	phosphate phosphatase	2641	putative protein
2611	hypothetical protein	2642	xanthine dehydrogenase
2612	putative flavonol 3-O-	2643	xanthine dehydrogenase - like
	glucosyltransferase	proteir	n
2613	60S ribosomal protein L30	2644	receptor protein kinase (IRK1),
2614	putative auxin-induced		putative
proteir	ı	2645	dehydrin-like protein
2615	putative nonspecific lipid-	2646	unknown protein
	transfer protein precursor	2647	aldehyde dehydrogenase homolog,
2616	AtRer1A		putative
2617	putative aquaporin	2648	Ran binding protein (AtRanBP1b)
	(tonoplast intrinsic protein	2649	putative squamosa-promoter
	gamma)		binding protein
2618	hypothetical protein	2650	putative protein
2619	putative alanine acetyl	2651	kinesin like protein
	transferase	2652	putative cellulose synthase
2620	putative NADP-dependent	2653	calmodulin (cam2)
	glyceraldehyde-3-	2654	fibrillarin - like protein
	phosphate dehydrogenase	2655	putative transmembrane protein
2621	putative DNA binding		G5p
proteir	1	2656	putative peroxidase
2622	putative cystathionine	2657	putative SNF1-related protein
	gamma-synthase		kinase
2623	unknown protein	2658	glutathione S-transferase, putative
2624	malate oxidoreductase	2659	unknown protein
	(malic enzyme)	2660	hypothetical protein
2625	unknown protein	2661	putative protein
2626	cyclic nucleotide-gated	2662	phosphatidylinositol-4-phosphate
	cation channel		5-kinase isolog
2627	glyoxalase II, putative	2663	putative tyrosine decarboxylase
2628	putative trypsin inhibitor	2664	unknown protein
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2665	SGP1 monomeric G-protein
	(emb CAB54517.1)
2666	putative serine
	carboxypeptidase II
2667	putative L5 ribosomal
protei	n
2668	putative glucosyltransferase
2669	flavonoid 3,5-hydroxylase
	like protein
2670	putative protein
2671	putative protein
2672	putative Fe(II)/ascorbate
	oxidase
2673	putative anthocyanin 5-
	aromatic acyltransferase
2674	casein kinase I
2675	putative 2,3-
	bisphosphoglycerate-
	independent
	phosphoglycerate mutase
2676	putative glutathione S-
	transferase TSI-1
2677	ATP-dependent RNA
helica	
2678	putative cytochrome P450
2679	putative WD-40 repeat
protei	
2680	No function assigned by
TIGR	
2681	No function assigned by
TIGR	
2682	putative protein
2683	putative extensin
2684	nodulin-26 - like protein
2685	RNA helicase
2686	(emb CAA09212.1)
2686	predicted protein of unknown function
2687	
2007	putative berberine bridge enzyme
2688	thioredoxin, putative
2689	putative serine
2009	carboxypeptidase I
2690	cytochrome P450-like
protei	
protei	u

2691	putative pyrophosphate-dependent
	phosphofructokinase alpha subunit
2692	putative flavonol
	glucosyltransferase
2693	peroxidase ATP20a
	(emb CAA67338.1)
2694	TOPP8 serine/threonine protein
	phosphatase type one
2695	auxin regulated protein IAA18,
	putative
2696	putative WRKY-type DNA binding
	protein
2697	putative glucan synthase
2698	squalene monooxygenase
2699	putative proline-rich protein
2700	G2484-1 protein
2701	heat shock protein 70 like protein
2702	unknown protein
2703	unknown protein
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TABLE 2
ABIOTIC STRESS RESPONSIVE GENE REGULATORY SEQUENCES

SEO RE	GULATORY	SEQ	REGULATORY	SEQ	REGULATORY
ID NO:	REGION	ID NO:	REGION	ID NO:	REGION
1	2704	51	2753	101	2802
2	2705	52	2754	102	2803
3	2706	53	2755	103	2804
4	2707	54	2756	104	2805
5	2708	55	2757	105	2806
6	2709	56	2758	106	2807
7	2710	57	2759	107	2808
8	2711	58	2760	107	2809
9	2712	59	2761	108	
10	2712	60	2762	110	2810
11	2714	61	2762		2811
12	2715	62		111	2812
13		63	2764	112	2813
	2716 2717		2765	113	2814
14		64	2766	114	2815
15	2718	65	2767	115	2816
16	2719	66	2768	116	2817
17	2720	67	2769	117	2818
18	2721	68	2770	118	2819
19	2722	69	NONE	119	2820
20	2723	70	2771	120	2821
21	2724	71	2772	121	2822
22	2725	72	2773	122	2823
23	2726	73	2774	123	2824
24	2727	74	2775	124	2825
25	2728	75	2776	125	2826
26	2729	76	2777	126	2827
27	2730	77	2778	127	2828
28	2731	78	2779	128	2829
29	2732	79	2780	129	2830
30	2733	80	2781	130	2831
31	2734	81	2782	131	2832
32	2735	82	2783	132	2833
33	2736	83	2784	133	2834
34	2737	84	2785	134	2835
35	2738	85	2786	135	2836
36	2739	86	2787	136	2837
37	2740	87	2788	137	2838
38	2741	88	2789	138	2839
39	2742	89	2790	139	2840
40	2743	90	2791	140	2841
41	2744	91	2792	141	2842
42	2745	92	2793	142	2843
43	NONE	93	2794	143	2844
44	2746	94	2795	144	NONE
45	2747	95	2796	145	2845
46	2748	96	2797	146	2846
47	2749	97	2798	147	2847
48	2750	98	2799	148	2848
49	2751	99	2800	149	2849
50	2752	100	2801	150	2850
- 0	2,52	100	2001	150	2000

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151	2851	205	2905	259	2959
152	2852	206	2906	260	2960
153	2853	207	2907	261	2961
154	2854	208	2908	262	2962
155	2855	209	2909	263	2963
156	2856	210	2910	264	2964
157	2857	211	2911	265	2965
158	2858	212	2912	266	2966
159	2859	213	2912	267	2967
160	2860	214	2914	268	2968
161	2861	215	2915	269	2969
162	2862	216	2916	270	2970
163	2863	217	2917	271	2970
164	2864	218	2918	272	2972
165	2865	219	2919	272	2972
166	2866	220	2920	274	2974
167	2867	221	2921	275	2975
168	2868	222	2921	276	2975 2976
169	2869	223	2922	277	2976 2977
170	2870	224	2923	278	2977
171	2871	225	2924	279	
171	2872		2925		2979
		226		280	2980
173	2873	227	2927	281	2981
174	2874	228	2928	282	2982
175	2875	229	2929 2930	283	2983
176 177	2876	230 231	2931	284 285	2984
	2877 2878		2931		2985
178		232 233	2932	286 287	2986 2987
179	2879		2933 2934		
180 181	2880 2881	234 235	2934	288 289	2988 2989
182	2882	236	2936	290	2990
183	2883	230	2936	290	
184	2884	238	2938	292	2991 2992
185	2885	238	2938	292	2992
186	2886	240	2939 2940	294	2993 2994
187	2887	240	2940 2941	295	2994
188	2888	241	2941	296	2995 2996
189	2889	242	2942	297	2997
190	2890	244	2943	298	2998
190	2891	244	2944	298	2999
191	2892	246	2945	300	3000
192	2893	247	2946	301	3001
194	2894	248	2947	302	3001
194	2895	249	2948	303	3002
195	2896	250	2950	304	NONE
196	2897	250	2951	305	3004
197	2898	251	2951	306	3004
198	2898 2899	252	2952 2953	306	3005 3006
200	2900	253	2954	308	3000
200	2900 2901	254 255	2954 2955	308	3007
201	2902	256	2956	310	3009
202	2902	250 257	2956 2957	311	3010
203	2903	257	2957 2958	311	3010
20 4	∠904	230	2930	31∠	3011

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313	3012	367	3066	421	3120
314	3013	368	3067	422	3121
315	3014	369	3068	423	3122
316	3015	370	3069	424	3123
317	3016	371	3070	425	3124
318	3017	372	3071	426	3125
319	3018	373	3072	427	3126
320	3019	374	3073	428	3127
321	3020	375	3074	429	3128
322	3021	376	3075	430	3129
323	3022	377	3076	431	3130
324	3023	378	3077	432	3131
325	3024	379	3078	433	3132
326	3025	380	3079	434	3133
327	3026	381	3080	435	3134
328	3027	382	3081	436	3135
329	3028	383	3082	437	3136
330	3029	384	3083	438	3137
331	3030	385	3084	439	3138
332	3031	386	3085	440	3139
333	3032	387	3086	441	3140
334	3033	388	3087	442	3141
335	3034	389	3088	443	3142
336	3035	390	3089	444	3143
337	3036	391	3090	445	3144
338	3037	392	3091	446	3145
339	3038	393	3092	447	3146
340	3039	394	3093	448	3147
341	3040	395	3094	449	3148
342	3041	396	3095	450	3149
343	3042	397	3096	451	3150
344	3043	398	3097	452	3151
345	3044	399	3098	453	3152
346	3045	400	3099	454	3153
347	3046	401	3100	455	3154
348	3047	402	3101	456	3155
349	3048	403	3102	457	3156
350	3049	404	3103	458	3157
351	3050	405	3104	459	3158
352	3051	406	3105	460	3159
353	3052	407	3106	461	3160
354	3053	408	3107	462	3161
355	3054	409	3108	463	3162
356	3055	410	3109	464	3163
357	3056	411	3110	465	3164
358	3057	412	3111	466	3165
359	3058	413	3112	467	3166
360	3059	414	3113	468	3167
361	3060	415	3114	469	3168
362	3061	416	3115	470	3169
363	3062	417	3116	471	3170
364	3063	418	3117	472	3171
365	3064	419	3118	473	3172
366	3065	420	3119	474	3173

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475	3174	529	3228	583	3282
476	3175	530	3229	584	3283
477	3176	531	3230	585	3284
478	3177	532	3231	586	3285
479	3178	533	3232	587	3286
480	3179	534	3233	588	3287
481	3180	535	3234	589	3288
482	3181	536	3235	590	3289
483	3182	537	3236	591	3290
484	3183	538	3237	592	3291
485	3184	539	3238	593	3292
486	3185	540	3239	594	3293
487	3186	541	3240	595	3294
488	3187	542	3241	596	3295
489	3188	543	3242	597	3296
490	3189	544	3243	598	3297
491	3190	545	3244	599	3298
492	3191	546	3245	600	3299
493	3192	547	3246	601	3300
494	3193	548	3247	602	3301
495	3194	549	3248	603	3302
496	3195	550	3249	604	3303
497	3196	551	3250	605	3304
498	3197	552	3251	606	3305
499	3198	553	3252	607	3306
500	3199	554	3253	608	3307
501	3200	555	3254	609	3308
502	3201	556	3255	610	3309
503	3202	557	3256	611	3310
504	3203	558	3257	612	3311
505	3204	559	3258	613	3312
506	3205	560	3259	614	3313
507	3206	561	3260	615	3314
508	3207	562	3261	616	3315
509	3208	563	3262	617	3316
510	3209	564	3263	618	3317
511	3210	565	3264	619	3318
512	3211	566	3265	620	3319
513	3212	567	3266	621	3320
514	3213	568	3267	622	3321
515	3214	569	3268	623	3322
516	3215	570	3269	624	3323
517	3216	571	3270	625	3324
518	3217	572	3271	626	3325
519	3218	573	3272	627	3326
520	3219	574	3273	628	3327
521	3220	575	3274	629	3328
522	3221	576	3275	630	3329
523	3222	577	3276	631	3330
524	3223	578	3277	632	3331
525	3224	579	3278	633	3332
526	3225	580	3279	634	3333
527	3226	581	3280	635	3334
528	3227	582	3281	636	3335

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637	3336	691	3390	745	3444
638	3337	692	3391	746	3445
639	3338	693	3392	747	3446
640	3339	694	3393	748	3447
641	3340	695	3394	749	3448
642	3341	696	3395	750	3449
643	3342	697	3396	751	3450
644	3343	698	3397	752	3451
645	3344	699	3398	753	3452
646	3345	700	3399	754	3453
647	3346	701	3400	755	3454
648	3347	702	3401	756	3455
649	3348	703	3402	757	3456
650	3349	704	3403	758	3457
651	3350	705	3404	759	3458
652	3351	706	3405	760	3459
653	3352	707	3406	761	3460
654	3353	708	3407	762	3461
655	3354	709	3408	763	3462
656	3355	710	3409	764	3463
657	3356	711	3410	765	3464
658	3357	712	3411	766	3465
659	3358	713	3412	767	3466
660	3359	714	3413	768	3467
661	3360	715	3414	769	3468
662	3361	716	3415	770	3469
663	3362	717	3416	771	3470
664	3363	718	3417	772	3471
665	3364	719	3418	773	3472
666	3365	720	3419	774	3473
667	3366	721	3420	775	3474
668	3367	722	3421	776	3475
669	3368	723	3422	777	3476
670	3369	724	3423	778	3477
671	3370	725	3424	779	3478
672	3371	726	3425	780	3479
673	3372	727	3426	781	3480
674	3373	728	3427	782	3481
675	3374	729	3428	783	3482
676	3375	730	3429	784	3483
677	3376	731	3430	785	3484
678	3377	732	3431	786	3485
679	3378	733	3432	787	3486
680	3379	734	3433	788	3487
681	3380	735	3434	789	3488
682	3381	736	3435	790	3489
683	3382	737	3436	791	3490
684	3383	738	3437	792	3491
685	3384	739	3438	793	3492
686	3385	740	3439	794	3493
687	3386	741	3440	795	3494
688	3387	742	3441	796	3495
689	3388	743	3442	797	3496
690	3389	744	3443	798	3497

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799	3498	853	3552	907	3603
800	3499	854	3553	908	3604
801	3500	855	3554	909	3605
802	3501	856	3555	910	3606
803	3502	857	3556	911	3607
804	3503	858	3557	912	3608
805	3504	859	3558	913	3609
806	3505	860	3559	914	3610
807	3506	861	3560	915	3611
808	3507	862	3561	916	3612
809	3508	863	3562	917	3613
810	3509	864	3563	918	3614
811	3510	865	3564	919	3615
812	3511	866	3565	920	3616
813	3512	867	3566	921	3617
814	3513	868	3567	922	3618
815	3514	869	3568	923	3619
816	3515	870	3569	924	3620
817	3516	871	3570	925	3621
818	3517	872	3571	926	3622
819	3518	873	3572	927	3623
820	3519	874	3573	928	3624
821	3520	875	3574	929	3625
822	3521	876	3575	930	3626
823	3522	877	3576	931	3627
824	3523	878	3577	932	3628
825	3524	879	3578	933	3629
826	3525	880	3579	934	3630
827	3526	881	3580	935	NONE
828	3527	882	3581	936	3631
829	3528	883	3582	937	3632
830	3529	884	3583	938	3633
831	3530	885	3584	939	3634
832	3531	886	3585	940	3635
833	3532	887	NONE	941	3636
834	3533	888	3586	942	3637
835	3534	889	3587	943	3638
836	3535	890	3588	944	3639
837	3536	891	3589	945	3640
838	3537	892	3590	946	3641
839	3538	893	3591	947	3642
840	3539	894	NONE	948	3643
841	3540	895	NONE	949	3644
842	3541	896	3592	950	3645
843	3542	897	3593	951	3646
844	3543	898	3594	952	3647
845	3544	899	3595	953	3648
846	3545	900	3596	954	3649
847	3546	901	3597	955	3650
848	3547	902	3598	956	3651
849	3548	903	3599	957	3652
850	3549	904	3600	958	3653
851	3550	905	3601	959	3654
852	3551	906	3602	960	3655

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962	3657	1016	3711	1070	3765
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967	3662	1021	3716	1075	3770
968	3663	1022	3717	1076	3771
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970	3665	1024	3719	1078	3773
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972	3667	1026	3721	1080	3775
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974	3669	1028	3723	1082	3777
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979	3674	1033	3728	1087	NONE
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981	3676	1035	3730	1089	3783
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994	3689	1048	3743	1102	3796
995	3690	1049	3744	1103	3797
996	3691	1050	3745	1104	3798
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1001	3696	1055	3750	1109	3803
1002	3697	1056	3751	1110	3804
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1006	3701	1060	3755	1114	3808
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1144	3838	1198	3892	1252	3946
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1327	4021	1381	4074	1435	4127
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1329	4023	1383	4076	1437	4129
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1783	4472	1837	4524	1891	4578
1784	NONE	1838	4525	1892	4579
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1803	NONE	1857	4544	1911	4596
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2127	4811	2181	4864	2235	4918
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2131	4815	2185	4868	2239	4922
2132	4816	2186	4869	2240	4923
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2134	4818	2188	4871	2242	4925
2135	4819	2189	4872	2243	4926
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2138	4822	2192	4875	2246	4929
2139	4823	2193	4876	2247	4930
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2144	4828	2198	4881	2252	4934
2145	4829	2199	4882	2253	4935
2146	4830	2200	4883	2254	4936
2147	4831	2201	4884	2255	4937
2148	4832	2202	4885	2256	4938

			, ,		
2257	4939	2311	4993	2365	5046
2258	4940	2312	4994	2366	5047
2259	4941	2313	4995	2367	5048
2260	4942	2314	4996	2368	5049
2261	4943	2315	4997	2369	5050
2262	4944	2316	4998	2370	5051
2263	4945	2317	4999	2371	NONE
2264	4946	2318	5000	2372	5052
2265	4947	2319	5001	2373	5053
2266	4948	2320	5002	2374	5054
2267	4949	2321	5002	2375	5055
2268	4950	2322	5004	2376	5056
2269	4951	2323	5005	2377	5057
2270	4952	2324	5005	2378	5058
2271	4953	2325	5007	2379	5059
2272	4954	2326	5007	2380	5060
2273	4955	2327	5009	2381	5061
2274	4956	2328	5010	2382	5062
2274	4957	2329	5011	2383	5063
2276	4958	2330	5012	2384	5064
2276	4958 4959	2331	5012	2385	5065
	4959 4960	2332	5014	2386	5066
2278					5067
2279	4961	2333	5015	2387	
2280	4962	2334	5016	2388 2389	5068 5069
2281	4963	2335	5017	2390	5070
2282	4964	2336 2337	5018 5019	2390	5070
2283	4965				
2284	4966	2338	5020	2392	5072
2285	4967	2339	5021	2393	5073
2286	4968	2340	NONE	2394	5074 5075
2287	4969	2341	5022	2395	
2288	4970	2342	5023	2396 2397	5076 5077
2289	4971	2343	5024		
2290	4972	2344	5025	2398	5078
2291	4973	2345	5026	2399	5079
2292	4974	2346	5027	2400	5080
2293	4975	2347	5028	2401	5081
2294	4976	2348	5029	2402	5082
2295	4977	2349	5030	2403	5083
2296	4978	2350	5031	2404	5084
2297	4979	2351	5032	2405	5085
2298	4980	2352	5033	2406	5086
2299	4981	2353	5034	2407	5087
2300	4982	2354	5035	2408	5088
2301	4983	2355	5036	2409	5089
2302	4984	2356	5037	2410	5090
2303	4985	2357	5038	2411	5091
2304	4986	2358	5039	2412	5092
2305	4987	2359	5040	2413	5093
2306	4988	2360	5041	2414	5094
2307	4989	2361	5042	2415	5095
2308	4990	2362	5043	2416	5096
2309	4991	2363	5044	2417	5097
2310	4992	2364	5045	2418	5098

2419	5099	2473	5151	2527	5205
2420	5100	2474	5152	2528	5206
2421	5101	2475	5153	2529	5207
2422	5102	2476	5154	2530	5208
2423	5103	2477	5155	2531	5209
2424	5104	2478	5156	2532	5210
2425	5105	2479	5157	2533	5211
2426	5106	2480	5158	2534	5212
2427	5107	2481	5159	2535	5213
2428	5108	2482	5160	2536	5214
2429	5109	2483	5161	2537	5215
2430	5110	2484	5162	2538	5216
2431	5111	2485	5163	2539	5217
2432	5112	2486	5164	2540	5218
2433	5113	2487	5165	2541	5219
2434	5114	2488	5166	2542	5219
2435	5115	2489	5167	2543	5220
2435	5116	2490	5168	2544	5221
2430	5117	2490	5169	2545 2545	5222 5223
2437	5118	2491	5170	2546	5223
2438	5119	2492			
			5171	2547	5225
2440	5120	2494	5172	2548	5226
2441	5121	2495	5173	2549	5227
2442	5122	2496	5174	2550	5228
2443	NONE	2497	5175	2551	5229
2444	5123	2498	5176	2552	5230
2445	5124	2499	5177	2553	5231
2446	5125	2500	5178	2554	5232
2447	5126	2501	5179	2555	5233
2448	5127	2502	5180	2556	5234
2449	5128	2503	5181	2557	5235
2450	5129	2504	5182	2558	5236
2451	5130	2505	5183	2559	5237
2452	5131	2506	5184	2560	5238
2453	5132	2507	5185	2561	5239
2454	5133	2508	5186	2562	5240
2455	5134	2509	5187	2563	5241
2456	5135	2510	5188	2564	5242
2457	5136	2511	5189	2565	5243
2458	5137	2512	5190	2566	5244
2459	5138	2513	5191	2567	5245
2460	5139	2514	5192	2568	5246
2461	5140	2515	5193	2569	5247
2462	5141	2516	5194	2570	5248
2463	5142	2517	5195	2571	5249
2464	5143	2518	5196	2572	5250
2465	5144	2519	5197	2573	5251
2466	5145	2520	5198	2574	5252
2467	5146	2521	5199	2575	5253
2468	5147	2522	5200	2576	5254
2469	NONE	2523	5201	2577	5255
2470	5148	2524	5202	2578	5256
2471	5149	2525	5203	2579	5257
2472	5150	2526	5204	2580	5258

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2581	5259	2635	5312
2582	5260	2636	5313
2583	5261	2637	5314
2584	5262	2638	5315
2585	5263	2639	5316
2586	5264	2640	5317
2587	5265	2641	5318
2588	5266	2642	5319
2589	5267	2643	5320
2590	5268	2644	5321
2591	5269	2645	5322
2592	5270	2646	5323
2593	5271	2647	5324
2594	5272	2648	5325
2595	5273	2649	5326
2596	5274	2650	5327
2597	5275	2651	5328
2598	5276	2652	5329
2599	NONE	2653	5330
2600	5277	2654	5331
2601	5278	2655	5332
2602	5279	2656	5333
2602	5280	2657	5334
2604	5281	2658	5335
2605	5282	2659	5336
2606	5283	2660	5337
2606	5284	2661	5338
2608	5285	2662	5339
2609	5286	2663	5340
2610	5287	2664	5341
2611	5288	2665	5342
2612	5289	2666	5343
2613	5290	2667	5344
2614	5291	2668	5345
2615	5292	2669	5346
2616	5293	2670	5347
2617	5294	2671	5348
2618	5295	2672	5349
2619	5296	2673	5350
2620	5297	2674	5351
2621	5298	2675	5352
2622	5299	2676	5353
2623	5300	2677	5354
2624	5301	2678	5355
2625	5302	2679	5356
2626	5303	2680	5357
2627	5304	2681	NONE
2628	5305	2682	5358
2629	5306	2683	5359
2630	5307	2684	5360
2631	5308	2685	5361
2632	5309	2686	5362
2633	5310	2687	5363
2634	5311	2688	5364

2689	5365
2690	5366
2691	5367
2692	5368
2693	5369
2694	5370
2695	5371
2696	5372
2697	5373
2698	5374
2699	5375
2700	5376
2701	5377
2702	5378
2703	5379

TABLE 3

COLD RESPONSIVE SEQUENCES

	PER COMPANY	aro.	A PER O APPROVA	aro	AFFYMETRIX
	AFFYMETRIX	SEQ	AFFYMETRIX	SEQ ID NO:	
ID NO:	ID NO:	ID NO:	ID NO:	98	ID NO:
1	11991_G_AT	50	12269_S_AT	98	12550_S_AT
2	11992_AT	51	12270_AT	99	17103_S_AT 12552_AT
3	11997_AT	52	12284_AT		
4	11998_AT	53	12287_S_AT	100	12555_S_AT
5	12001_AT		17570_G_AT	101	12576_S_AT
6	12006_S_AT	54	12293_AT	102	12581_S_AT
7	12007_AT	55	12294_S_AT		16645_S_AT
8	12009_AT	56	12300_AT	103	12587_AT
9	12018_AT	57	12307_AT	104	12597_AT
10	12022_AT	58	12312_AT	105	12602_AT
11	12026_AT	59	12315_AT	106	12610_AT
12	12031_AT	60	12324_I_AT	107	12631_AT
13	12047_AT	61	12331_S_AT	108	12646_AT
14	12051_AT	62	12336_AT	109	12649_AT
15	12052_AT	63	12344_AT	110	12650_AT
16	12053_AT	64	12348_AT	111	12653_AT
17	12060 AT	65	12353_AT	112	12661_AT
18	12072 AT	66	12359 S AT	113	12666_AT
19	12074 AT	67	12372_AT	114	12674_AT
20	12102 AT	68	12374 I AT	115	12675_S_AT
21	12112 AT		12726_F_AT	116	12678_I_AT
22	12117 AT	69	12390 AT	117	12681_S_AT
23	12125 AT	70	12395 S AT	118	12688_AT
24	12130_AT	71	12405 AT	119	12702 AT
25	12143 AT	72	12408 AT	120	12705 F_AT
26	12145 S AT	73	12410 G AT	121	12736 F AT
27	12149 AT	74	12419 AT	122	12737_F_AT
28	12156 AT	75	12427_AT	123	12758 AT
29	12163 AT	76	12431 AT	124	12760 G AT
30	12166 I AT	77	12436 AT	125	12762 R AT
31	12167 AT	78	12438 AT	126	12764 F AT
32	12169 I AT	79	12443 S AT	127	12766 AT
33	12175 AT	80	12447 AT		15115 F_AT
34	12175_AT	81	12450 S AT	128	12767 AT
35	12170_AT 12179_AT	82	12452 AT	129	12768 AT
36	12175_AT 12187_AT	83	12474 AT	130	12772 AT
30	15920 I AT	84	12477 AT	131	12773 AT
37	12195 AT	85	12491 AT	132	12776 AT
38	12195_AT 12196 AT	86	12497 AT	133	12788_AT
39	12198 AT	87	12500_S_AT	134	12793 AT
	12200 AT	88	12503 AT	135	12794 AT
40 41	12200_AT 12202_AT	89	12515 AT	136	12802 AT
42	12202_AT 12214 G AT	90	12516 S AT	137	12809 G AT
		91	12510_S_AT	138	12812 AT
43	12219_AT	92	12526 AT	139	12812_AT 12815_AT
44	12224_AT	92		140	12815_AT
45	12226_AT	93	12527_AT	141	12818 AT
46	12233_AT	94 95	12532_AT 12534 G AT	142	12824 S AT
47	12240_AT	95 96		142	12828 S AT
48	12253_G_AT		12544_AT	143	
49	12256_AT	97	12549_S_AT	144	12842_S_AT

145	12846_S_AT	194	13086_R_AT	238	13285_S_AT
146	12858 AT	195	13087 AT	239	13288 S AT
147	12860 S AT	196	13090 AT		17043 S AT
148	12861 S AT	197	13092 S AT	240	13292 S AT
149	12881 S AT	.,,	16950_S_AT	241	13296_S_AT
147	17600 S AT	198	13098 AT	242	13297 S AT
150				243	
150	12889_S_AT	199	13100_AT	243	13299_S_AT
151	12901_S_AT	200	13103_AT		15166_S_AT
152	12902_AT	201	13105_AT	244	13332_AT
153	12904_S_AT	202	13107_S_AT	245	13347_AT
154	12905_S_AT	203	13108_AT	246	13351_AT
155	12908 S AT	204	13109_AT	247	13352_AT
156	12910 S AT	205	13114 AT	248	13355_AT
	16385 S AT	206	13118 F AT	249	13404 AT
157	12914 S AT	207	13119 AT	250	13422 AT
	15783 S AT	208	13120 AT	251	13459 AT
	17645 S AT	209	13123 AT	252	13460 AT
158	12916 S AT	210	13128 AT	253	13461 S AT
159	12910_3_AT	211	13133 S AT	254	13467 AT
		211	17430 S AT	255	13488 AT
160	12926_S_AT	212		256	
161	12927_S_AT	212	13135_S_AT		13523_S_AT
162	12931_S_AT	213	13139_AT	257	13529_AT
163	12937_R_AT	214	13140_AT	258	13539_I_AT
164	12941_G_AT	215	13143_AT		14631_S_AT
165	12942_AT	216	13151_G_AT	259	13541_AT
166	12947_AT	217	13160_AT	260	13542_AT
167	12949 AT	218	13161_AT	261	13545_S_AT
168	12953 AT	219	13162_AT	262	13552_AT
169	12956 I AT	220	13165 AT	263	13556 I AT
170	12959 AT	221	13166 AT	264	13561 AT
171	12966 S AT	222	13167 AT	265	13563 S AT
172	12975 AT	223	13179 AT	266	13567 AT
173	12983 AT	224	13181 AT	267	13568 AT
174	12984 AT	225	13185 AT	268	13571 AT
175	12987_S_AT	226	13193 S AT	269	13575 AT
		227	13213 S AT	270	13576 AT
176	12994_S_AT	221		271	13583_AT
177	13002_AT	220	16004_S_AT	271	
178	13009_I_AT	228	13219_S_AT		13598_AT
179	13011_AT		20288_G_AT	273	13601_AT
180	13018_AT	229	13220_S_AT	274	13604_AT
181	13023_AT		13221_AT	275	13613_AT
182	13024_AT		18929_S_AT	276	13616_S_AT
183	13034_S_AT	230	13233_AT		16544_S_AT
184	13046 G_AT		14301_S_AT	277	13617_AT
185	13048 S_AT	231	13243 R AT	278	13618_S_AT
	13495 S AT	232	13254 S AT	279	13619_AT
186	13054 AT	233	13260 S AT	280	13621 G AT
187	13067 S_AT		15660 S AT	281	13623 R AT
188	13068 AT	234	13273 S AT	282	13629 S AT
189	13073 S AT		16105 S AT	283	13631 AT
190	13078 S AT	235	13274_S_AT	284	13635 AT
191	13079 AT	230	17077 S AT	285	13646 AT
191	13081 S AT	236	13276 S AT	286	13650 AT
192	13081_S_A1 13083 AT	237	13278 F AT	287	13653_AT
193	12002 W1	231	132/0_F_A1	201	12022_W1

288	13655 AT	332	13989 AT	383	14393 AT
289	13656 AT		20674 S AT	384	14421 AT
290	13657_AT	333	14010 AT	385	14436 AT
291	13666 S AT	334	14013 AT	386	14448 AT
	17083 S AT	335	14014 AT	387	14450 AT
292	13667_S_AT	336	14019_AT	388	14454_AT
293	13669 S AT	337	14021 R AT	389	14459 AT
	17074_S_AT	338	14025 S AT	390	14478 AT
294	13670 S AT		18909 S AT	391	14482 AT
	15206 S AT	339	14027 AT	392	14485 AT
295	13671 S AT	340	14030 AT	393	14492 S AT
	16805 S AT	341	14044 AT	394	14505 AT
296	13678 S AT	342	14048 AT	395	14510 AT
297	13688 S AT	343	14056 AT	396	14511 AT
298	13690 S AT	344	14057 AT	397	14517 AT
	16065 S AT	345	14058 AT	398	14519 AT
299	13691 S AT	346	14059 AT	399	14525 S AT
2,,	16117 S AT	347	14061 AT	400	14527 AT
300	13692 S AT	348	14068 S AT	401	14534 S AT
500	16118 S AT	349	14072 AT	402	14538 R AT
301	13700 AT	350	14073 AT	403	14554 AT
302	13704 S AT	351	14074 AT	404	14558 AT
303	13714 AT	352	14084 AT	405	14559 S AT
304	13715 AT	353	14095 S AT	406	14566 AT
305	13724 AT	354	14100 AT	407	14572 AT
306	13748 AT	355	14101 AT	408	14579 AT
307	13759 AT	356	14103 AT	409	14587 AT
308	13767 AT	357	14105 AT	410	14591 AT
309	13785 AT	358	14106 AT	411	14595 AT
310	13803 AT	359	14121 AT	412	14602 AT
311	13850 I AT	360	14129 S AT	413	14603 AT
312	13876 AT	361	14133_S_AT	414	14605 AT
313	13880 S AT	362	14143 AT	415	14620 S AT
314	13883 AT	363	14145 AT	416	14626_S_AT
315	13887 S AT	364	14148 AT	417	14630 S AT
316	13895 AT	365	14186 AT		16559 S AT
317	13904 S AT	366	14194 AT	418	14637 S AT
	18722 S AT	367	14196 AT		17122 S AT
318	13906 S AT	368	14223_AT	419	14642_F_AT
319	13908 S AT	369	14234 AT	420	14650_S_AT
	18597 AT	370	14236_AT		15150_S_AT
320	13923 AT	371	14251_F_AT	421	14654_S_AT
321	13927_AT	372	14252_F_AT	422	14667_S_AT
322	13932_AT	373	14270_AT		18299_S_AT
323	13935_AT	374	14298_G_AT	423	14669_S_AT
324	13940_AT		17581_G_AT		16136_S_AT
325	13949_S_AT	375	14303_S_AT	424	14672_S_AT
326	13954_G_AT	376	14312_AT	425	14679_S_AT
327	13971_S_AT	377	14316_AT	426	14682_I_AT
328	13973_AT	378	14339_AT	427	14689_AT
329	13983_AT	379	14366_AT	428	14697_G_AT
330	13985_S_AT	380	14369_AT		16902_AT
331	13987_S_AT	381	14388_AT	429	14701_S_AT
	18738_F_AT	382	14392_G_AT		14734_S_AT

430	14703_AT	483	15130_S_AT	534	15489_AT
431	14711_S_AT	484	15131_S_AT	535	15490_AT
432	14712_S_AT	485	15132_S_AT	536	15503_AT
	20530_S_AT		17585_S_AT	537	15505_AT
433	14713_S_AT	486	15139_S_AT	538	15510_R_AT
434	14715 S AT	487	15143 S AT	539	15512 AT
435	14728 S AT	488	15146 S AT	540	15514 AT
436	14731 S AT	489	15159 S AT	541	15515 R AT
437	14781 AT		15160 S AT	542	15517 S AT
438	14797 S AT	490	15162 S AT	543	15518 AT
439	14800 AT	491	15167 S AT	544	15529 AT
440	14809_AT	492	15171_S_AT	545	15534_F_AT
441	14843 AT	493	15174 F AT	546	15538 AT
442	14847_AT	494	15178 S AT	547	15541 AT
443	14872 AT	495	15185 S AT	548	15543 AT
444	14886 AT		18023 S AT	549	15544 AT
445	14896 AT	496	15188_S_AT	550	15551_AT
446	14900 AT	497	15193 S AT	551	15574 S AT
447	14908 AT	498	15196 S AT	552	15576 S AT
448	14912 AT	499	15197 S AT	553	15577 S AT
449	14914 AT	500	15201 F AT	554	15578 S AT
450	14942 AT	501	15213 S AT	555	15583 S AT
451	14945 AT	502	15243 AT	556	15588 S AT
452	14955 AT	503	15256 AT	557	15595 S AT
453	14957 S AT	504	15270 AT	558	15600 S AT
454	14958 AT	505	15319 AT	559	15602 F_AT
455	14965 AT	506	15325 AT	560	15608_S_AT
456	14974 AT	507	15337 AT	561	15613 S AT
457	14980 AT	508	15341 AT	562	15616 S AT
458	14981 AT	509	15343_AT	563	15618 S AT
459	14984 S AT	510	15348 AT	564	15620_S_AT
460	14995_AT	511	15350_AT	565	15627_S_AT
461	15004_AT	512	15355_S_AT	566	15634_S_AT
462	15009_AT	513	15367_AT		16125_S_AT
463	15010 AT	514	15372_AT		18046_S_AT
464	15024_AT	515	15379_AT	567	15637_S_AT
465	15026_AT	516	15381_AT	568	15639_S_AT
466	15036_R_AT	517	15383_AT	569	15642_S_AT
467	15054_AT	518	15384_AT	570	15643_S_AT
468	15056_AT	519	15385_AT	571	15651_F_AT
469	15057_AT	520	15387_AT	572	15652_S_AT
470	15066_AT	521	15410_AT	573	15665_S_AT
471	15073_AT	522	15417_S_AT	574	15667_S_AT
472	15081_AT	523	15422_AT		18610_S_AT
473	15083_AT	524	15423_AT	575	15668_S_AT
474	15091_AT	525	15431_AT	576	15671_S_AT
475	15097_S_AT	526	15433_AT	577	15675_S_AT
476	15101_S_AT	527	15452_AT	578	15679_S_AT
477	15102_S_AT	528	15464_AT	579	15685_S_AT
478	15107_S_AT	529	15468_AT	580	15687_F_AT
479	15112_S_AT	530	15471_AT	581	15688_S_AT
480	15116_F_AT	531	15472_AT	582	15689_S_AT
481	15118_S_AT	532	15475_S_AT	583	15692_S_AT
482	15122_S_AT	533	15485_AT	584	15694_S_AT

585	15712_S_AT	634	16089_S_AT	686	16496_S_AT
586	15808_AT	635	16090_S_AT	687	16499_AT
587	15845_AT	636	16102 S AT	688	16510 AT
588	15848_AT	637	16103_S_AT	689	16511_AT
589	15850_AT	638	16108 S AT	690	16512 S AT
	20406_G_AT	639	16112_S_AT		18085_R_AT
590	15858 AT	640	16134 S AT	691	16514 AT
591	15862 AT	641	16137 S AT	692	16516 AT
592	15868 AT	642	16138 S AT	693	16517 AT
593	15878 AT	643	16140 S AT	694	16526 AT
594	15894_AT	644	16143 S AT	695	16528_AT
595	15900 AT	645	16145 S AT	696	16531 S AT
596	15901 AT	646	16148 S AT	697	16535 S AT
597	15902 AT	647	16151 S AT	698	16537 S AT
598	15912 AT	648	16155 S AT	699	16538 S AT
599	15913 AT	649	16158 F AT	700	16543 S AT
600	15928 AT	650	16160 F AT	701	16550 S AT
601	15940 AT	651	16162 S AT	702	16554 S AT
602	15941 AT	652	16168 S AT	703	16567 S AT
603	15945 AT	653	16169 S AT	704	16571 S AT
604	15948 S AT	654	16171 S AT	705	16576 F AT
605	15956 AT	655	16172 S AT	706	16577 S AT
606	15960 AT	656	16184 AT	707	16579 S AT
	16466 S AT	657	16192 AT	708	16580 S AT
607	15976 AT	658	16222 AT	709	16583 S AT
608	15978 AT	659	16242 AT	710	16584 S AT
609	15986 S AT	660	16244 AT		18706 S AT
610	15990 AT	661	16250 AT	711	16593 S AT
611	16009 S AT	662	16286_AT	712	16595_S_AT
612	16015 AT	663	16288 AT	713	16598 S AT
613	16019 AT	664	16294 S AT	714	16604 S AT
614	16024 AT	665	16296_AT	715	16605_S_AT
615	16034 AT	666	16297 AT	716	16610_S_AT
616	16036 I AT	667	16325_AT	717	16611_S_AT
	18729_AT	668	16346_S_AT	718	16614_S_AT
617	16039 S AT	669	16357_AT	719	16617_S_AT
618	16040_AT	670	16380_AT	720	16618_S_AT
619	16042_S_AT	671	16382_AT	721	16620_S_AT
620	16047_AT	672	16393_S_AT	722	16621_S_AT
621	16049_S_AT	673	16402_S_AT	723	16631_S_AT
622	16051_S_AT	674	16411_S_AT	724	16634_S_AT
623	16055_S_AT	675	16442_S_AT	725	16635_S_AT
624	16059_S_AT	676	16446_AT	726	16636_S_AT
625	16062_S_AT	677	16448_G_AT	727	16639_S_AT
626	16066_S_AT	678	16453_S_AT	728	16640_S_AT
627	16069_S_AT	679	16457_S_AT	729	16650_S_AT
628	16074_S_AT	680	16465_AT	730	16652_S_AT
629	16076_S_AT		16916_S_AT	731	16654_AT
630	16077_S_AT	681	16470_S_AT	732	16672_AT
	17579_S_AT		18735_S_AT	733	16673_AT
631	16079_S_AT	682	16481_S_AT	734	16687_S_AT
632	16084_S_AT	683	16486_AT	735	16747_AT
	17998_S_AT	684	16487_AT	736	16753_AT
633	16087_S_AT	685	16488_AT	737	16768_AT

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738	16777 AT	790	17123 S AT	843	17562 AT
739	16784 AT	791	17129 S AT	844	17564 S AT
740	16807 AT	792	17132 AT	011	19361 S AT
741	16811 AT	793	17166 AT	845	17565 S AT
742	16845 AT	794	17206 AT	846	17568 AT
743	16894 AT	795	17207 AT	847	17573 AT
744	16899 AT	796	17215 AT	848	17577 G AT
745	16911 AT	797	17237 AT	849	17578 AT
746	16920 AT	798	17247 AT	850	17596 AT
747	16921 AT	799	17254 AT	851	17627 AT
748	16924_S_AT	800	17286 AT	852	17631 AT
749	16926 S AT	801	17288 S AT	853	17632 AT
750	16931 S AT	802	17292 AT	854	17672 AT
751	16934 S AT	803	17300 AT	855	17675 AT
752	16937 AT	804	17303 S AT	856	17677 AT
753	16938 AT	805	17318 AT	857	17732 AT
754	16942 AT	806	17319 AT	858	17743 AT
755	16943 S AT	807	17322 AT	859	17748 AT
133	18231 AT	808	17322_AT	860	17748_AT
756	16949 S AT	809	17332 S AT	861	17823 S AT
757	16952 S AT	810	17374 AT	862	17841 AT
758	16956 AT	811	17374_AT 17381 AT	863	17841_A1 17849 S AT
759	16962 S AT	812	17388 AT	864	17849_S_AT
760	16965 S AT	813	17392 S AT	865	17852_G_A1 17857_AT
761	16970 S AT	814	17405 AT	866	17865 AT
/01	18010 S AT	815	17405_AT	867	17882 AT
762	16977 AT	816	17415_A1 17418 S AT	868	17885_AT
763	16977_AT 16984 AT	817	17418_3_A1 17420 AT	869	17900 S AT
764	16996 S AT	818	17423 S AT	870	17900_S_AT
765	16997 AT	819	17426 AT	871	17911 AT
766	17000 AT	820	17427 AT	872	17916 AT
767	17005_AT	821	17429 S AT	873	17917 S AT
768	17010_S_AT	822	17431 AT	874	17918 AT
769	17010_S_AT	823	17439 G AT	875	17921 S AT
770	17031 S AT	824	17457 AT	876	17922 AT
771	17033_S_AT	825	17458_AT	877	17926 S AT
772	17053_5_AT	826	17462 S AT	878	17933 AT
773	17055_S_AT	827	17463 AT	879	17935_AT
774	17063 S AT	828	17465 AT	880	17956 I AT
775	17068 S AT	829	17466 S AT	881	17966 AT
776	17070 S AT	830	17475 AT	882	17967 AT
777	17075 S AT	831	17479 AT	883	17970 I AT
778	17084 S AT	832	17482 S AT	884	17978 S AT
779	17087 S AT	833	17495 S AT	001	20635 S AT
780	17092 S AT	834	17508 S AT	885	17986 S AT
781	17095 S AT	835	17522 S AT	886	17993 AT
782	17096 S AT	836	17523 S AT	887	18001 AT
783	17102 S AT	837	17537 S AT	888	18003 AT
784	17102_S_AT	838	17538 S AT	889	18004 AT
785	17109_S_AT	839	17539 S AT	890	18005_AT
786	17110_S_AT	840	17546 S AT	891	18029 G AT
787	17113 S AT	,	18694 S AT		18030 I AT
788	17115 S AT	841	17557 S AT	892	18040 S AT
789	17116 S AT	842	17560 S AT	893	18045 AT

894	18064_R_AT	947	18580 AT	1001	18889 AT
895	18065 R AT	948	18581 AT	1002	18892 S AT
896	18074 AT	949	18584 AT	1003	18901 AT
897	18076 S AT	950	18587 S AT	1004	18911 AT
898	18077_AT	951	18588 AT	1005	18917 I AT
899	18081 AT	952	18591 AT	1006	18939 AT
900	18154 S AT	953	18592 S AT	1007	18947 I AT
	18365 S AT	954	18600 AT	1008	18950 AT
901	18165 AT	955	18601 S AT	1009	18951 S AT
902	18174 AT	956	18607 S AT	1010	18954 AT
903	18176 AT	957	18611 AT	1011	18956 AT
904	18194 I AT	958	18616 AT	1012	18959 AT
905	18197 AT	959	18622 G AT	1013	18966 AT
906	18198 AT	960	18623 AT	1014	18974 AT
907	18213 AT	961	18628 AT	1015	18976 AT
908	18219 AT	962	18631 AT	1016	18980 AT
909	18221 AT	963	18635 AT	1017	18989 S AT
910	18222 AT	964	18636 AT	1018	18994 AT
911	18226 S AT	965	18638 AT	1019	19030 AT
912	18232 AT	966	18652 AT	1020	19039 AT
913	18237 AT	967	18657 AT	1021	19049 AT
914	18241 AT	968	18659 AT	1022	19083 AT
915	18257 AT	969	18660 S AT	1023	19115 AT
916	18258 S AT	970	18667 AT	1024	19117 S AT
917	18269 S AT	971	18675 AT	1025	19122 AT
918	18274 S AT	972	18684 AT	1026	19125 S AT
919	18275_AT	973	18686_S_AT	1027	19127_AT
920	18278_AT	974	18688 S AT	1028	19130_AT
921	18282_AT	975	18693_S_AT	1029	19144_AT
922	18283_AT	976	18698_S_AT	1030	19157_S_AT
923	18290_AT	977	18705_AT	1031	19178_AT
924	18291_AT	978	18707_AT	1032	19190_G_AT
925	18306_AT	979	18708_AT	1033	19198_AT
926	18316_AT	980	18726_S_AT	1034	19202_AT
927	18317_AT	981	18727_AT	1035	19209_S_AT
928	18327_S_AT	982	18732_I_AT	1036	19211_AT
929	18337_S_AT	983	18736_AT	1037	19218_AT
930	18339_AT	984	18750_F_AT	1038	19222_AT
931	18347_S_AT	985	18754_AT	1039	19226_G_AT
932	18383_AT	986	18778_AT	1040	19229_AT
933	18390_AT	987	18806_S_AT	1041	19230_AT
934	18439_S_AT	988	18823_S_AT	1042	19232_S_AT
935	18465_S_AT	989	18829_AT	1043	19285_AT
936	18487_AT	990	18835_AT	1044	19326_AT
937	18508_S_AT	991	18844_AT	1045	19332_AT
938	18512_AT	992	18859_AT	1046	19346_AT
939	18543_AT	993	18864_AT	1047	19347_AT
940	18544_AT	994	18866_AT	1048	19362_AT
941	18552_AT	995	18880_AT	1049	19363_AT
942	18555_AT	996	18883_G_AT	1050	19364_AT
943	18556_AT	997	18885_AT	1051	19367_AT
944	18561_AT	998	18886_AT	1052	19373_AT
945	18567_AT	999	18887_AT	1053	19381_AT
946	18573_AT	1000	18888_AT	1054	19382_AT

1055 19384 AT						
1058 19416_AT	1055	19384_AT	1109	19833 S AT	1163	20093 I AT
1058	1056	19401 AT	1110	19834 AT	1164	20099 AT
1058	1057	19406 AT	1111	19836 AT	1165	20100 AT
1059	1058	19413 AT	1112	19841 AT		
1060	1059		1113			
1061 1949 AT						
1062						
1063						
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1068 19470 AT						
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1070						
1071 19513						
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1073 19562_AT						
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1075 19567						
1076 1958						
1077 19589 S. AT 1131 19913 AT 1185 20255 AT 1177 19589 S. AT 1132 19920 S. AT 1186 20257 AT 11079 19606 AT 1133 19932 AT 1187 20262 AT 11080 19623 AT 1134 19939 AT 1188 20275 AT 1180 19624 AT 1135 19945 AT 1189 20278 S. AT 1081 19624 AT 1135 19945 AT 1189 20278 S. AT 1082 19627 S. AT 1136 19947 AT 1190 20282 S. AT 1083 19636 AT 1137 19951 AT 1191 20284 AT 1083 19635 AT 1138 19956 AT 1192 20293 AT 1083 19655 AT 1138 19956 AT 1192 20293 AT 1085 19657 S. AT 1140 19963 AT 1194 20312 S. AT 1086 19657 S. AT 1140 19963 AT 1194 20312 S. AT 1088 19660 AT 1142 19970 S. AT 1196 20330 S. AT 1089 19665 S. AT 1144 19971 AT 1197 20331 AT 1090 19667 AT 1144 19971 AT 1197 20331 AT 1090 19667 AT 1146 19990 AT 1200 20355 AT 1092 19677 AT 1146 19990 AT 1200 20355 AT 1093 19686 AT 1147 19990 AT 1200 20355 AT 1094 19689 AT 1148 20003 S. AT 1202 20363 AT 1094 19689 AT 1148 20003 S. AT 1202 20363 AT 1095 19690 S. AT 1149 20009 S. AT 1202 20363 AT 1096 19695 S. AT 1149 20009 S. AT 1202 20363 AT 1096 19695 S. AT 1150 20013 AT 1204 20378 G. AT 1097 19698 AT 1151 20018 AT 1204 20378 G. AT 1097 19698 AT 1151 20018 AT 1204 20378 G. AT 1097 19698 AT 1151 20018 AT 1206 20384 AT 1101 19726 S. AT 1155 20044 S. AT 1209 20395 AT 1101 19726 S. AT 1155 20044 S. AT 1204 20399 AT 1104 19759 AT 1156 20048 AT 1211 20409 G. AT 1104 19759 AT 1156 20048 AT 1212 20413 AT 1204 20378 AT 1104 19759 AT 1156 20058 AT 1212 20440 G. AT 1104 19782 AT 1156 20048 AT 1212 20440 G. AT 1106 19803 S. AT 1166 20067 AT 1214 20449 AT 1106 19803 S. AT 1166 20067 AT 1214 20449 AT 1106 19803 S. AT 1166 20067 AT 1214 20449 AT 1106 19803 S. AT 1166 2006						
1078 19595 S.AT 1132 19920 S.AT 1186 20257 AT 1187 20262 AT 1188 19606 AT 1131 19932 AT 1187 20262 AT 1188 19625 AT 1188 20275 AT 1188 20275 AT 1181 19624 AT 1135 19945 AT 1189 20278 S.AT 1081 19627 S.AT 1136 19947 AT 1190 20282 S.AT 1083 19636 AT 1137 19951 AT 1191 20284 AT 1084 19652 AT 1138 19956 AT 1192 20293 AT 1085 19655 AT 1138 19956 AT 1192 20293 AT 1085 19657 S.AT 1139 19962 AT 1193 20294 AT 1086 19657 S.AT 1140 19963 AT 1194 20312 S.AT 1087 19658 AT 1141 19969 AT 1195 20315 L.AT 1088 19655 S.AT 1141 19969 AT 1195 20315 L.AT 1088 19665 S.AT 1141 19970 S.AT 1196 20330 S.AT 1090 19667 AT 1144 19972 AT 1198 20350 S.AT 1092 19671 AT 1146 19990 AT 1199 20355 AT 1092 19677 AT 1146 19990 AT 1200 20355 AT 1093 19686 AT 1147 19996 AT 1200 20355 AT 1094 19689 AT 1148 20003 S.AT 1202 20363 AT 1095 19690 S.AT 1148 20003 S.AT 1202 20363 AT 1095 19690 S.AT 1149 20009 S.AT 1202 20363 AT 1096 19695 AT 1150 20013 AT 1204 20338 G.AT 1097 19698 AT 1151 20013 AT 1204 20338 G.AT 1097 19698 AT 1151 20013 AT 1204 20338 G.AT 1098 19700 S.AT 1152 20024 S.AT 1205 20338 AT 1099 19708 AT 1152 20024 S.AT 1206 20338 AT 1100 1917 AT 1154 20003 AT 1204 20338 G.AT 1101 19726 S.AT 1155 20047 AT 1209 20396 AT 1101 19726 S.AT 1156 20048 AT 1212 20409 G.AT 1104 19759 AT 1158 20051 AT 1214 20409 G.AT 1104 19759 AT 1158 20051 AT 1214 20449 AT 1106 19803 S.AT 1160 20067 AT 1214 20449 AT 1106 19803 S.AT 1160 20068 AT 1215 20440 AT 1216 20440 AT 1216 20440 AT 12068 AT 1215 20440 AT 12068 AT 12068 AT 1						
1079 19606 AT						
1080 19623_AT						
1081 19624 AT						
1082 19627 S. AT 1136 19947 AT 1190 20282 S. AT 1083 19536 AT 1137 19951 AT 1191 20284 AT 1084 19652 AT 1138 19956 AT 1192 20293 AT 1085 19655 AT 1139 19962 AT 1193 20294 AT 1086 19657 S. AT 1140 19963 AT 1194 20312 S. AT 1087 19658 AT 1141 19969 AT 1195 20315 F. AT 1088 19660 AT 1142 19971 AT 1197 20331 AT 1090 19665 S. AT 1144 19971 AT 1197 20331 AT 1090 19667 AT 1144 19972 AT 1198 20350 S. AT 1091 19671 AT 1145 19981 AT 1199 20354 S. AT 1092 19677 AT 1146 19990 AT 1200 20355 AT 1093 19686 AT 1144 19990 AT 1202 20363 AT 1094 19689 AT						
1083 19636 AT						
1084 19652 AT						
1085 19655.AT 1139 19962.AT 1193 20294.AT 1086 19657.S.AT 1140 19963.AT 1194 20312.S.AT 1087 19658.AT 1141 19969.AT 1195 20315.LAT 1088 19660.AT 1142 19970.SAT 1196 20330.S.AT 1089 19665.S.AT 1144 19971.AT 1197 20331.AT 1090 19667.AT 1144 19972.AT 1198 20350.S.AT 1091 19671.AT 1146 19990.AT 1200 20355.AT 1092 19677.AT 1146 19990.AT 1200 20355.AT 1093 19686.AT 1147 19996.AT 1201 20360.AT 1094 19689.AT 1148 20003.S.AT 1202 20363.AT 1095 19690.S.AT 1149 20009.S.AT 1202 20369.S.AT 1096 19695.AT 1151 20013.AT 1204 20378.G.AT 1098 19900.S.AT						
1086 19657. S. AT 1140 19963. AT 1194 20312. S. AT 1087 19658. AT 1141 19969. AT 1195 20315. LAT 1088 19660. AT 1142 19970. S. AT 1196 20330. S. AT 1089 19665. S. AT 1143 19971. AT 1197 20331. AT 1090 19667. AT 1144 19972. AT 1198 20350. S. AT 1091 19671. AT 1145 19981. AT 1199 20354. S. AT 1092 19677. AT 1146 19990. AT 1200 20355. AT 1093 19686. AT 1147 19996. AT 1201 20360. AT 1094 19689. AT 1148 20003. S. AT 1202 20360. AT 1095 19690. S. AT 1149 20009. S. AT 1202 20369. S. AT 1096 19695. AT 1150 20013. AT 1204 20378. G. AT 1097 19698. AT 1151 20013. AT 1204 20378. G. AT						
1087 19658 AT						
1088 19660 AT						
1089 19665 S AT						
1090 19667 AT						
1091 1967 AT						
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1993 19686 AT						
1094 19689 AT						
1095 19690 S AT 1149 20009 S AT 1203 20366 S AT 1096 19695 AT 1150 20013 AT 1204 20378 G AT 1097 19698 AT 1151 20018 AT 1205 20383 AT 1098 19700 S AT 1152 20024 S AT 1206 20384 AT 1109 19708 AT 1153 20027 AT 1207 20387 AT 1100 19717 AT 1154 20045 AT 1208 20393 AT 1101 19726 S AT 1155 20047 AT 1209 20396 AT 1102 19744 AT 1156 20048 AT 1210 20399 AT 1103 19752 S AT 1157 20050 AT 1211 20409 G AT 1104 19759 AT 1158 20051 AT 1212 20412 S AT 1105 1982 AT 1159 20058 AT 1212 20412 S AT 1105 1983 S AT 1160 20067 AT 1214 20439 AT 1107 19828 AT <						
1096						
1997 19968_AT 1151 20018_AT 1205 20383_AT 1098 19700_S_AT 1152 20024_S_AT 1206 20384_AT 1099 19708_AT 1153 20027_AT 1207 20387_AT 1100 19717_AT 1154 20045_AT 1208 20393_AT 1101 19726_S_AT 1155 20047_AT 1209 20396_AT 1102 19744_AT 1156 20048_AT 1210 20399_AT 1103 19752_S_AT 1157 20050_AT 1211 20409_G_AT 1104 19759_AT 1158 20051_AT 1212 20412_S_AT 1105 19803_S_AT 1169 20058_AT 1213 20413_AT 1106 19803_S_AT 1160 20067_AT 1214 20439_AT 1107 19828_AT 1161 20068_AT 1215 20440_AT						
1098 19700 S AT 1152 20024 S AT 1206 20384 AT 1099 19708 AT 1153 20027 AT 1207 20387 AT 1100 19717 AT 1154 20045 AT 1208 20393 AT 1101 19726 S AT 1155 20047 AT 1209 20396 AT 1102 19744 AT 1156 20048 AT 1210 20399 AT 1103 19752 S AT 1157 20050 AT 1211 20409 G AT 1104 19759 AT 1158 20051 AT 1212 20412 S AT 1105 19782 AT 1159 20058 AT 1213 20413 AT 1106 19803 S AT 1160 20067 AT 1214 20439 AT 1107 19828 AT 1160 20068 AT 1215 20440 AT						
1099 19708_AT 1153 20027_AT 1207 20387_AT 1100 19717_AT 1154 20045_AT 1208 20393_AT 1101 19726_S_AT 1155 20047_AT 1209 20396_AT 1102 19744_AT 1156 20048_AT 1210 20399_AT 1103 19752_S_AT 1157 20050_AT 1211 20409_G_AT 1104 19759_AT 1158 20051_AT 1212 20412_S_AT 1105 19803_S_AT 1159 20058_AT 1213 20413_AT 1106 19803_S_AT 1160 20067_AT 1214 20439_AT 1107 19828_AT 1161 20068_AT 1215 20440_AT						
1100 19717_AT						
1101 19726 S. AT 1155 20047 AT 1209 20396 AT 1102 19744 AT 1156 20048 AT 1210 20399 AT 1103 19752 S. AT 1157 20050 AT 1211 20409 G. AT 1104 19759 AT 1158 20051 AT 1212 20412 S. AT 1105 19782 AT 1159 20058 AT 1213 20413 AT 1106 19803 S. AT 1160 20067 AT 1214 20439 AT 1107 19828 AT 1161 20068 AT 1215 20440 AT 1215 20440 AT 1216 20490 AT 1216 20490 AT 1217 20490 AT 1218 20490						
1102 19744_AT 1156 20048_AT 1210 20399_AT 1103 19752_SAT 1157 20050_AT 1211 20409_G_AT 1104 19759_AT 1158 20051_AT 1212 20412_S_AT 1105 19782_AT 1159 20058_AT 1213 20413_AT 1106 19803_S_AT 1160 20067_AT 1214 20439_AT 1107 19828_AT 1161 20068_AT 1215 20440_AT						
1103 19752_S AT 1157 20050_AT 1211 20409_G AT 1104 19759_AT 1158 20051_AT 1212 20412_S AT 1105 19782_AT 1159 20058_AT 1213 20413_AT 1106 19803_S_AT 1160 20067_AT 1214 20439_AT 1107 19828_AT 1161 20068_AT 1215 20440_AT						
1104 19759 AT 1158 20051 AT 1212 20412 S AT 1105 19782 AT 1159 20058 AT 1213 20413 AT 1106 19803 S AT 1160 20067 AT 1214 20439 AT 1107 19828 AT 1161 20068 AT 1215 20440 AT						
1105 19782_AT 1159 20058_AT 1213 20413_AT 1106 19803_S_AT 1160 20067_AT 1214 20439_AT 1107 19828_AT 1161 20068_AT 1215 20440_AT						
1106 19803_S_AT 1160 20067_AT 1214 20439_AT 1107 19828_AT 1161 20068_AT 1215 20440_AT						
1107 19828_AT 1161 20068_AT 1215 20440_AT						
1108 19831_1_A1 1162 20069_AT 1216 20444_AT						
	1108	19831_1_AT	1162	20069_A1	1216	20444_AT

1217	20445 AT
1218	20449_AT
1219	20456_AT
1220	20462_AT
1221	20471_AT
1222	20474_AT
1223	20495 S AT
1224	20499 AT
1225	20501 AT
1226	20511 AT
1227	20515 S AT
1228	20516 AT
1229	20517 AT
1230	20518 AT
1231	20520_S_AT
1232	20536 S AT
1233	20538 S AT
1234	20539 S AT
1235	20558_AT
1236	20561 AT
1237	20567 AT
1238	20571 AT
1239	20582 S AT
1240	20586 I AT
1241	20590 AT
1242	20592 AT
1242	20592_AT
1244	20608 S AT
1245	20612 S AT
1246	20616 AT
1247	20620 G AT
1248	20637_AT
1249	20643 AT
1250	20649 AT
1251	20651 AT
1252	20654 S AT
1253	20670 AT
1254	20684 AT
1255	20685 AT
1256	20693 AT
1257	20701 S AT
1258	20704_AT
1259	20705 AT
1260	20715 AT
1261	20719 AT

168 TABLE 4: 2X UP IN COLD, ONLY

11997_at	12688_at	13274_s_at	14145_at	15083_at	15639_s_at
11998_at	12701_i_at	13278_f_at	14170_at	15084_at	15641_s_at
12018_at	12702_at	13279_s_at	14186_at	15096_at	15660_s_at
12031_at	12719_f_at	13285_s_at	14196_at	15101_s_at	15665_s_at
12047_at	12726_f_at	13288_s_at	14227 at	15105_s_at	15687 f at
12051 at	12736 f at	13292 s at	14234 at	15112 s at	15694_s_at
12053 at	12754 g at	13297_s_at	14250 r at	15115 f at	15712_s_at
12060 at	12762 r at	13299 s at	14270 at	15116_f_at	15783 s at
	12762_1_at	13332 at	14298 g at	151122 s at	15808 at
12072_at	_				_
12074_at	12767_at	13351_at	14303_s_at	15126_s_at	15837_at
12102_at	12768_at	13352_at	14312_at	15131_s_at	15850_at
12112_at	12773_at	13422_at	14339_at	15132_s_at	15862_at
12117_at	12788_at	13435_at	14388_at	15137_s_at	15868_at
12130_at	12802_at	13461_s_at	14393_at	15144_s_at	15878_at
12145_s_at	12860_s_at	13467_at	14511_at	15148_s_at	15901_at
12151_at	12861_s_at	13488_at	14525_s_at	15153_s_at	15912_at
12163_at	12879_s_at	13495_s_at	14527_at	15159_s_at	15920_i_at
12175_at	12891_at	13539_i_at	14534_s_at	15 1 60_s_at	15941_at
12187_at	12914_s_at	13542_at	14554_at	15166_s_at	15945_at
12195_at	12927_s_at	13575_at	14566_at	15174_f_at	15960_at
12219_at	12947_at	13577_s_at	14579_at	15197_s_at	15990_at
12256_at	12956_i_at	13617_at	14591_at	15270_at	16001_at
12269 s at	12966_s_at	13634_s_at	14595_at	15319_at	16009_s_at
12307 at	12974 at	13656_at	14600_at	15325 at	16010_s_at
12315_at	12987_s_at	13671_s_at	14631_s_at	15337 at	16034_at
12336 at	12994 s at	13691 s at	14635 s at	15341 at	16036 i at
12349_s_at	12998 at	13700 at	14679 s at	15343 at	16039 s at
12353 at	13002_at	13704_s at	14691_at	15355_s_at	16040 at
12359_s_at	13018_at	13709_s_at	14697_g_at	15367_at	16042_s_at
12390 at	13023 at	13715 at	14709 at	15379 at	16047 at
12395 s at	13046 g at	13785 at	14711 s at	15381 at	16049_s_at
12431 at	13054 at	13803_at	14728_s_at	15410_at	16051 s at
12436_at	13086_r_at	13812_s_at	14731_s_at	15417_s_at	16062_s_at
12443_s_at	13087_at	13825 s at	14797_s_at	15422 at	16079_s_at
12447 at	13100 at	13850 i at	14809 at	15433_at	16087_s_at
12452 at	13109 at	13904_s_at	14843 at	15451 at	16090_s_at
12477_at	13119 at	13908 s at	14847 at	15452 at	16117_s_at
12503_at	13120 at	13927_at	14872 at	15453_s_at	16118_s_at
12516 s at	13128 at	13971 s at	14886 at	15472 at	16137_s_at
12532 at	13134_s_at	13985_s_at	14896_at	15489 at	16155_s_at
12544 at	13140 at	14013_at	14897 at	15490 at	16162 s at
12561_at	13143_at	14019_at	14900_at	15503_at	16184_at
12602_at	13143_at	14021_r_at	14956_s_at	15510_r_at	16192_at
12610_at	13172_s_at	14028_at	14958_at	15517_s_at	16222_at
12631_at	13178_at	14048_at	14965_at	15518_at	16244_at
12647_s_at	13179_at	14058_at	14984_s_at	15544_at	16250_at
12650_at	13181_at	14059_at	15004_at	15588_s_at	16260_at
12656_at	13187_i_at	14064_at	15010_at	15600_s_at	16286_at
12674_at	13209_s_at	14073_at	15036_r_at	15605_s_at	16296_at
12675_s_at	13219_s_at	14105_at	15040_g_at	15613_s_at	16297_at
12676_s_at	13221_at	14106_at	15046_s_at	15614_s_at	16342_at
12681_s_at	13243_r_at	14126_s_at	15057_at	15616_s_at	16367_i_at
12686_s_at	13260_s_at	14140_at	15073_at	15633_s_at	16411_s_at

169 TABLE 4 (cont): 2X UP IN COLD, ONLY

	1/10//	3 4 (cont). 2/L	or in Cold,	ONLI	
16442_s_at	17077_s_at	17978_s_at	18885_at	19689_at	20412_s_at
16465_at	17102_s_at	17999_at	18887_at	19698_at	20413_at
16466_s_at	17109_s_at	18001_at	18888_at	19700_s_at	20432_at
16468_at	17113_s_at	18004_at	18889_at	19707_s_at	20433_at
16486_at	17123_s_at	18012_s_at	18901_at	19708_at	20456_at
16487_at	17128_s_at	18040_s_at	18907_s_at	19713_at	20462_at
16488_at	17129_s_at	18176_at	18917_i_at	19718_at	20471_at
16489_at	17132_at	18194_i_at	18939_at	19744_at	20511_at
16496_s_at	17166_at	18197_at	18947_i_at	19836_at	20515_s_at
16499_at	17206_at	18198_at	18949_at	19839_at	20517_at
16511_at	17237_at	18213_at	18954_at	19840_s_at	20518_at
16517_at	17300_at	18219_at	18959_at	19845_g_at	20529_at
16538_s_at	17319_at	18222_at	18974_at	19854_at	20536_s_at
16554 s at	17322 at	18231 at	18976_at	19855_at	20538_s_at
16571 s at	17332 s at	18232_at	18980_at	19860_at	20539_s_at
16576_f_at	17381_at	18241_at	18989_s_at	19866_at	20576_at
16595_s_at	17388_at	18269_s_at	19019_i_at	19871_at	20582_s_at
16605_s_at	17392_s_at	18272_at	19049_at	19875_s_at	20586_i_at
16610_s_at	17408 at	18282_at	19083_at	19879_s_at	20608_s_at
16620_s_at	17424 at	18298 at	19130_at	19881_at	20649_at
16621 s at	17429_s_at	18316_at	19156_s_at	19913_at	20651_at
16635_s_at	17457_at	18317_at	19178_at	19939_at	20684_at
16636_s_at	17458_at	18331_s_at	19190_g_at	19945_at	20685_at
16638_s_at	17466_s_at	18347_s_at	19199_at	19947_at	20699_at
16650_s_at	17477_s_at	18383_at	19202_at	19951_at	20705_at
16672_at	17482_s_at	18390_at	19209_s_at	19956_at	20715_at
16673_at	17538_s_at	18455_at	19211_at	19971_at	
16687_s_at	17546_s_at	18465_s_at	19218_at	19976_at	
16747_at	17562_at	18544_at	19229_at	19998_at	
16753_at	17581_g_at	18555_at	19322_at	20003_s_at	
16768_at	17627_at	18556_at	19326_at	20015_at	
16805_s_at	17631_at	18560_at	19359_s_at	20027_at	
16807_at	17632_at	18561_at	19367_at	20051_at	
16845_at	17645_s_at	18571_at	19384_at	20068_at	
16847_at	17672_at	18588_at	19389_at	20093_i_at	
16896_s_at	17675_at	18597_at	19397_at	20117_at	
16899_at	17677_at	18601_s_at	19406_at	20150_at	
16902_at	17693_at	18611_at	19426_s_at	20156_at	
16911_at	17732_at	18623_at	19441_s_at	20165_at	
16914_s_at	17743_at	18635_at	19442_at	20257_at	
16943_s_at	17748_at	18659_at	19470_at	20262_at	
16956_at	17775_at	18660_s_at	19489_s_at	20275_at	
16996_s_at	17782_at	18673_at	19562_at	20282_s_at	
17010_s_at	17841_at	18694_s_at	19577_at	20288_g_at	
17016_s_at 17032_s_at	17852_g_at	18705_at 18708_at	19589_s_at	20293_at	
	17900_s_at 17901_at		19597_s_at	20315_i_at	
17033_s_at 17043 s at	17901_at	18738_f_at 18750_f_at	19611_s_at 19624_at	20330_s_at 20360_at	
17043_s_at	17921_s_at	18778_at	19657_s_at	20363 at	
17050_s_at	17922_at	18829 at	19667 at	20369_s_at	
17068_s_at	17933_at	18835_at	19671_at	20384_at	
17071_s_at	17967_at	18866_at	19677_at	20393 at	
17075 s at	17970 i at	18875 s at	19686_at	20396_at	

170 TABLE 5: 2X UP COLD 3 HR, ONLY

19624 at 19657_s_at 19667_at 19845_g_at 19855 at 19866 at 19945 at 19951 at 19998_at 20003 s at 20015_at 20051 at 20093_i_at 20117_at 20288_g_at 20360 at

20369_s_at

20608_s_at 20651_at 20685_at 20705_at

20384_at 20462_at 20471_at 20515_s_at 20538_s_at 20576_at

	IADEE	. 2.1 01 0022 0	,
12117_at	13671_s_at	15453_s_at	17237_at
12145_s_at	13691_s_at	15489_at	17319_at
12151_at	13785_at	15518_at	17392_s_at
12163_at	13803_at	15588_s_at	17429_s_at
12187_at	13825_s_at	15613_s_at	17477_s_at
12256_at	13904_s_at	15614_s_at	17538_s_at
12315_at	14013_at	15616_s_at	17581_g_at
12349_s_at	14021_r_at	15639_s_at	17627_at
12353_at	14028_at	15641_s_at	17672_at
12359_s_at	14064_at	15660_s_at	17693_at
12544 at	14126_s_at	15687_f_at	17782_at
12602 at	14145_at	15694_s_at	17841_at
12610 at	14170_at	15862 at	17900_s_at
12676_s_at	14196_at	15868_at	17933 at
12686 s at	14250_r_at	15878 at	17978 s at
12701_i_at	14298 g_at	15901 at	18001 at
12702 at	14303 s at	16034_at	18012_s_at
12719 f at	14339 at	16039_s_at	18198 at
12736 f at	14527_at	16040 at	18219 at
12754 g at	14534_s_at	16042 s at	18241 at
12766 at	14554 at	16047 at	18269 s_at
12767 at	14595_at	16062_s_at	18272 at
12767_at	14635 s at	16087 s at	18282 at
12773_at	14679_s_at	16117 s at	18298 at
12775_at 12788 at	14691 at	16117_5_at	18383 at
12700_at	14697_g_at	16162 s at	18556 at
12891_at	14709 at	16184 at	18588 at
12947 at	14705_at	16222 at	18601 s at
12947_at	14720_3_at	16250 at	18611_at
12974 at	14896 at	16411 s at	18694_s_at
12994 s at	14965 at	16442 s at	18708 at
13002_at	14984 s at	16465 at	18738 f at
13100 at	15046 s at	16486 at	18778 at
13140 at	15083_at	16488 at	18829 at
13167 at	15096 at	16489 at	18835 at
13172 s at	15105 s at	16517 at	18866 at
13179 at	15115 f at	16571 s at	18875_s_at
13187 i at	15116_f_at	16605_s_at	18888 at
13219 s at	15122 s_at	16610 s at	18907 s at
13260 s at	15126 s at	16620_s_at	18917 i at
13278 f at	15131 s at	16636 s at	18939 at
13279 s at	15132 s at	16650_s_at	18974 at
13285_s_at	15137_s_at	16805 s at	19190_g_at
13288 s at	15153 s at	16845 at	19199_at
13292_s_at	15159 s at	16899 at	19202_at
13297 s at	15160 s at	16914_s_at	19211 at
13351_at	15197_s_at	16943 s at	19384 at
13351_at	15355 s at	16996 s at	19406 at
13435 at	15355_s_at 15379 at	17010 s at	19426 s at
13435_at	15379_at 15417 s at	17010_s_at	19442 at
13488_at	15417_s_at	17043_s_at	19470 at
13488_at	15422_at 15451 at	17066_s_at	19577_at
	15451_at	17109_s_at	19597_s_at
13656_at	1040Z_dt	11 120_5_al	15551_5_dl

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TABLE 6: 2X DOWN COLD, ONLY

11001 a at	124E0 a at	12881 s at	13151 g at	13621 g_at	14056_at
11991_g_at	12450_s_at				
11992_at	12474_at	12889_s_at	13160_at	13623_r_at	14057_at
12001_at	12491_at	12901_s_at	13161_at	13629_s_at	14061_at
12006_s_at	12497_at	12902_at	13162_at	13631_at	14067_at
12007_at	12500_s_at	12904_s_at	13165_at	13635_at	14068_s_at
12009_at	12515_at	12905_s_at	13166_at	13646_at	14072_at
12022_at	12521_at	12908_s_at	13185_at	13650_at	14074_at
12023_s_at	12523_at	12910_s_at	13193_s_at	13652_at	14075_at
12026_at	12526_at	12916_s_at	13211_s_at	13653_at	14083_at
12037_at	12527_at	12923_s_at	13213_s_at	13655_at	14084_at
12052_at	12534_g_at	12926_s_at	13219_s_at	13657_at	14089_at
12125_at	12549_s_at	12931_s_at	13233_at	13666_s_at	14095_s_at
12143_at	12550_s_at	12937_r_at	13236_s_at	13667_s_at	14096_at
12149_at	12552_at	12941_g_at	13239_s_at	13669_s_at	14100_at
12156_at	12555_s_at	12942 at	13241_s_at	13670_s_at	14101_at
12166_i_at	12556 at	12949_at	13254 s at	13672 s at	14103_at
12167 at	12575 s at	12953 at	13266 s at	13678 s at	14121 at
12169 i at	12576 s at	12958 at	13273_s_at	13679_s_at	14129 s at
12176 at	12581_s_at	12959 at	13275 f at	13688 s at	14133 s at
12179_at	12587 at	12966 s at	13276 s at	13690_s_at	14143_at
12196_at	12597_at	12975 at	13278 f at	13691 s at	14148 at
12198 at	12606 at	12983 at	13280 s at	13692 s at	14162_at
12200_at	12609_at	12984_at	13285_s_at	13714 at	14194 at
12200_at	12646 at	13002 at	13296 s at	13724_at	14208_at
12212 at	12649 at	13009_i_at	13347 at	13748 at	14217 at
12212_at	12653 at	13011 at	13355 at	13751 at	14223_at
	12661 at	13014_at	13361_at	13759 at	14235 at
12224_at	12666 at	13024_at	13404 at	13767_at	14236_at
12226_at 12233_at	12678_i_at	13034_s_at	13406_at	13789 at	14251 f at
		13041 s at	13459 at	13876 at	14252 f at
12240_at	12705_f_at 12736_f_at		13460 at	13880_s_at	14285 at
12253_g_at		13048_s_at	13464_at	13883 at	14301 s at
12270_at	12737_f_at	13067_s_at		13887_s_at	14301_s_at
12278_at	12758_at	13068_at	13523_s_at	13895_at	14366 at
12284_at	12760_g_at	13073_s_at	13529_at 13541_at	13906 s at	14369 at
12287_s_at	12764_f_at	13078_s_at	_		14392_g_at
12293_at	12765_at	13079_at	13545_s_at	13919_at 13923_at	14421 at
12294_s_at	12772_at	13081_s_at	13550_at		
12300_at	12776_at	13083_at	13552_at	13932_at	14431_at
12312_at	12784_at	13090_at	13556_i_at	13935_at	14436_at
12315_at	12793_at	13092_s_at	13561_at	13940_at	14448_at
12324_i_at	12794_at	13098_at	13563_s_at	13949_s_at	14450_at
12331_s_at	12795_at	13103_at	13567_at	13954_g_at	14454_at
12344_at	12809_g_at	13105_at	13568_at	13973_at	14459_at
12348_at	12812_at	13107_s_at	13571_at	13983_at	14478_at
12353_at	12815_at	13108_at	13576_at	13989_at	14482_at
12372_at	12816_at	13114_at	13583_at	14010_at	14485_at
12374_i_at	12818_at	13118_f_at	13598_at	14014_at	14492_s_at
12405_at	12824_s_at	13123_at	13601_at	14015_s_at	14505_at
12408_at	12828_s_at	13124_at	13604_at	14016_s_at	14510_at
12410_g_at	12842_s_at	13133_s_at	13613_at	14025_s_at	14517_at
12419_at	12846_s_at	13135_s_at	13616_s_at	14027_at	14519_at
12427_at	12858_at	13139_at	13618_s_at	14030_at	14534_s_at
12438_at	12869_s_at	13146_s_at	13619_at	14044_at	14538_r_at

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TABLE 6 (cont): 2X DOWN COLD, ONLY

4.4EEQ of	15047 of	45540 of	45040 ot	10057 -4	10001 -4
14558_at	15047_at	15512_at	15940_at	16357_at	16894_at
14559_s_at	15054_at	15514_at	15948_s_at	16380_at	16899_at
14572_at	15056_at	15515_r_at	15956_at	16382_at	16920_at
14584_at	15058_s_at	15529_at	15976_at	16385_s_at	16921_at
14587_at	15063_at	15534_f_at	15978_at	16393_s_at	16924_s_at
14595_at	15066_at	15538_at	15986_s_at	16402_s_at	16926_s_at
14602_at	15081_at	15541_at	16004_s_at	16417_s_at	16931_s_at
14603 at	15091_at	15543 at	16015_at	16442 s at	16934 s at
14605 at	15097 s at	15551_at	16017 at	16446 at	16937 at
14620 s at	15102 s_at	15574_s_at	16019_at	16448 g at	16938_at
14626 s at	15107_s_at	15576 s at	16024_at	16453 s at	16942 at
14630 s at	15118 s at	15577 s at	16031 at	16457 s at	16949 s at
14637 s at	15127_s_at	15578_s_at	16055_s_at	16470 s at	16950_s_at
14640_s_at	15130_s_at	15581_s_at	16059_s_at	16481_s_at	16952_s_at
14642 f at	15132_s_at	15583 s at	16065_s_at	16510 at	16962_s_at
14650 s at	15133 s at	15591 s at	16066 s at	16512 s at	16965 s at
14654 s at	15139 s at	15595_s_at	16069 s at	16514 at	16970 s at
14667 s at	15143_s_at	15602 f at	16074_s_at	16516 at	16977 at
14668_s_at	15146_s_at	15606_s_at	16076_s_at	16523_s_at	16984_at
14669 s at	15150 s at	15608_s_at	16077 s at	16526 at	16989 at
				16528 at	16993 at
14672_s_at	15161_s_at	15616_s_at	16084_s_at	_	_
14673_s_at	15162_s_at	15618_s_at	16089_s_at	16531_s_at	16997_at
14675_s_at	15167_s_at	15620_s_at	16102_s_at	16535_s_at	17000_at
14679_s_at	15170_s_at	15627_s_at	16103_s_at	16537_s_at	17005_at
14681_g_at	15171_s_at	15634_s_at	16105_s_at	16543_s_at	17010_s_at
14682_i_at	15178_s_at	15637_s_at	16108_s_at	16544_s_at	17017_s_at
14689_at	15182_s_at	15642_s_at	16112_s_at	16550_s_at	17031_s_at
14701_s_at	15185_s_at	15643_s_at	16117_s_at	16559_s_at	17040_s_at
14703_at	15188_s_at	15646_s_at	16118_s_at	16567_s_at	17053_s_at
14712_s_at	15193_s_at	15651_f_at	16125_s_at	16577_s_at	17056_s_at
14713_s_at	15196_s_at	15652_s_at	16127_s_at	16579_s_at	17063_s_at
14715_s_at	15201_f_at	15667_s_at	16134_s_at	16580_s_at	17070_s_at
14734_s_at	15206_s_at	15668_s_at	16136_s_at	16583_s_at	17074_s_at
14781_at	15207_s_at	15670_s_at	16138_s_at	16584_s_at	17084_s_at
14800_at	15213_s_at	15671_s_at	16140_s_at	16593_s_at	17085_s_at
14856_s_at	15243_at	15675_s_at	16143_s_at	16598_s_at	17087_s_at
14882_at	15256_at	15679_s_at	16144_s_at	16603_s_at	17092_s_at
14908_at	15348_at	15685_s_at	16145_s_at	16604_s_at	17095_s_at
14912_at	15350_at	15688_s_at	16148_s_at	16611_s_at	17096_s_at
14914_at	15372_at	15689_s_at	16151_s_at	16614_s_at	17097_s_at
14924_at	15383_at	15692_s_at	16158_f_at	16617_s_at	17103_s_at
14942_at	15384_at	15775_at	16160_ f_ at	16618_s_at	17105_s_at
14945_at	15385_at	15776_at	16168_s_at	16620_s_at	17110_s_at
14955_at	15387_at	15845_at	16169_s_at	16631_s_at	17115_s_at
14957_s_at	15406_at	15848_at	16171_s_at	16634_s_at	17116_s_at
14974_at	15423_at	15858_at	16172_s_at	16639_s_at	17119_s_at
14980_at	15431_at	15866_s_at	16222_at	16640_s_at	17122_s_at
14981_at	15464_at	15894_at	16232_s_at	16652_s_at	17207_at
14995_at	15468_at	15900_at	16242_at	16654_at	17215_at
15009_at	15471_at	15901_at	16288_at	16777_at	17247_at
15018_at	15475_s_at	15902_at	16294_s_at	16784_at	17254_at
15024_at	15485_at	15913_at	16325_at	16811_at	17286_at
15026_at	15505_at	15928_at	16346_s_at	16893_at	17288_s_at
_	_	_		_	

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TABLE 6 (cont): 2X DOWN COLD, ONLY

17292_at	17910_at	18337_s_at	18823_s_at	19382_at	19897_s_at
17303 s at	17916 at	18339_at	18844_at	19401_at	19903_at
17305 at	17917_s_at	18365_s_at	18859_at	19402_at	19905_at
17318 at	17918_at	18402_at	18864_at	19406_at	19906_at
17323_at	17926_s_at	18439_s_at	18880_at	19413_at	19907_at
17374_at	17935 at	18487_at	18883_g_at	19416_at	19910_at
17405_at	17956_i_at	18508_s_at	18886_at	19429_at	19920_s_at
17415 at	17961 at	18512 at	18892_s_at	19432_s_at	19932_at
17418 s at	17966_at	18543 at	18909_s_at	19439_at	19951_at
17420 at	17978 s at	18552 at	18911_at	19448_s_at	19962_at
17423 s_at	17986_s_at	18567_at	18913_s_at	19454_at	19963_at
17426 at	17993 at	18573_at	18916_s_at	19462_s_at	19969_at
17427_at	17998 s at	18580_at	18921 g at	19464 at	19970_s_at
17430 s at	18003 at	18581 at	18950 at	19469_at	19972_at
17431 at	18005 at	18584 at	18951 s_at	19483_at	19981_at
17439 g at	18010 s at	18587_s_at	18956_at	19484 s at	19990 at
17442 i at	18013_r_at	18590 at	18966 at	19513 at	19996 at
17449_s_at	18023_s_at	18591_at	18972_at	19548 at	19999_s_at
17462 s at	18029_g_at	18592 s at	18994 at	19563 s at	20009_s_at
17463 at	18030 i at	18600 at	19030 at	19567 at	20013_at
17465_at	18045 at	18601 s at	19039 at	19581_at	20017 at
17475 at	18046 s at	18607_s_at	19068 i at	19595 s at	20018 at
17479_at	18059_i_at	18610 s at	19108 at	19606_at	20024 s at
17475_at	18064 r at	18611 at	19115_at	19623 at	20045 at
17493_s_at	18065 r at	18616_at	19117_s_at	19627_s_at	20047_at
17500_s_at	18074_at	18622 g at	19122_at	19636_at	20048 at
17523 s at	18076_s_at	18628 at	19125_s_at	19641 at	20050 at
17529_s_at	18077_at	18631_at	19127_at	19652 at	20051 at
17525_s_at	18078 at	18636 at	19135_at	19655 at	20058 at
17537_s_at	18081_at	18638_at	19144 at	19658_at	20067_at
17533_s_at	18083 r at	18652_at	19157_s_at	19660_at	20069_at
17555 s_at	18085 r at	18657_at	19158 at	19665 s at	20099 at
17555_s_at	18091_at	18667_at	19177 at	19667 at	20100_at
17560 s_at	18154 s at	18675_at	19192 at	19690 s at	20113 s_at
17564_s_at	18165 at	18684 at	19198 at	19695_at	20123 at
17565 s at	18174_at	18686 s at	19222_at	19717_at	20127 s at
17568_at	18221 at	18688_s_at	19226 g_at	19726_s_at	20129_at
	18226 s at	18693_s_at	19227 at	19752 s at	20133 i at
17570_g_at 17573_at	18230_at	18698 s_at	19230 at	19759_at	20152 at
17573_at	18237 at	18706 s at	19232 s_at	19782_at	20154 at
17577_g_at	18255 at	18707 at	19263_at	19789 s at	20173 at
17579_s_at	18257_at	18726 s at	19285 at	19803_s_at	20178 s_at
17575_s_at	18258 s at	18727 at	19332_at	19828 at	20183 at
17596_at	18274_s_at	18732_i_at	19346 at	19831 i at	20188_at
17600_at	18275 at	18735 s at	19347_at	19833_s_at	20189_at
17823 s at	18278_at	18736_at	19361_s_at	19834 at	20197 at
17840_s_at	18283 at	18738 f at	19362 at	19835 at	20200 at
17849 s at	18290 at	18747 f at	19363 at	19841 at	20210 g at
17857 at	18291_at	18754 at	19364 at	19867_at	20213_at
17865 at	18299 s at	18782 at	19365 s at	19870_s_at	20229_at
_	18300 at	18789_at	19373_at	19871 at	20232_s_at
17882_at 17885_at	18306_at	18806_s_at	19379_at	19872 at	20255_at
	18327_s_at	18814_at	19375_at	19876 at	20278 s at
17902 s_at	10021_5_al	100 14_at	10001_41	.50,0_40	

TABLE 6 (cont): 2X DOWN COLD, ONLY

```
20284 at
            20693 at
20288 g at
            20701_s_at
            20704 at
20294 at
20312 s at
            20707 s at
20331_at
            20719_at
20335_s_at
20350_s_at
20354 s at
20355 at
20369 s at
20378_g_at
20383 at
20385_s_at
20387_at
20399 at
20409 g_at
20420 at
20429_s_at
20439 at
20440 at
20444 at
20445 at
20449_at
20474 at
20480 s at
20495_s_at
20499_at
20501 at
20516_at
20520_s_at
20530 s at
20538 s at
20547_at
20558 at
20561 at
20567 at
20571 at
20590 at
20592_at
20594 at
20608_s_at
20612 s at
20616 at
20620 g at
20635 s at
20637_at
20643 at
20654 s at
20670_at
20674 s at
20684 at
20685 at
```

20689_s_at

TABLE 7

	SALINE STRESS RESPONSIVE SEQUENCES						
SEO AFFYMETRIX SEO AFFYMETRIX SEO AFFYMETRIX							
ID NO:	ID NO:	ID NO:	ID NO:	ID N	O: ID NO:		
2227	12011_S_AT	2275	13993 S AT	2324	15965 AT		
2228	12153 AT	2276	14000 AT	2325	15969 S AT		
2229	12180 AT	2277	14003 AT	2326	15975_S_AT		
2230	12186 AT	2278	14032 AT	2327	15995 S AT		
2231	12216 AT	2279	14043 AT	2328	15998 S AT		
2232	12265 AT	2280	14070 AT		18090 S AT		
2233	12335_AT	2281	14267 AT	2329	16028_AT		
2234	12449 S AT	2282	14269 AT	2330	16050_AT		
2235	12470 AT	2283	14418_AT	2331	16060_S_AT		
2236	12479 AT	2284	14427_AT	2332	16067_S_AT		
2237	12487_AT	2285	14501_AT	2333	16072_S_AT		
2238	12493_G_AT	2286	14544_AT	2334	16088_F_AT		
2239	12562_AT	2287	14546_S_AT	2335	16273_AT		
2240	12685_AT	2288	14570_AT	2336	16314_AT		
2241	12704_F_AT	2289	14596_AT	2337			
2242	12709_F_AT	2290	14729_S_AT	2338			
2243	12734_F_AT	2291	14874_AT	2339			
2244	12739_S_AT	2292	14888_AT	2340			
2245	12750_S_AT	2293	14951_AT	2341	16455_AT		
2246	12761_S_AT	2294	14952_AT	2342			
2247	12813_AT	2295	14959_AT	2343			
2248	12845_S_AT	2296	14979_AT	2344			
2249	12946_AT	2297	15006_AT	2345			
2250	13003_S_AT	2298	15042_AT	2346			
2251	13052_S_AT	2299	15049_AT	2347 2348			
2252	13094_AT	2300 2301	15062_AT 15108 S AT	2349			
2253	13142_AT	2301	15147 S AT	2350			
2254	13172_S_AT	2302	15175 S AT	2351			
2255	17880_S_AT 13198 I AT	2304	15175_S_AT	2352			
2256	13209 S AT	2305	15186 S AT	2353			
2230	16165 S AT	2505	18696 S AT	2354			
2257	13229 S AT	2306	15192 S AT	2355			
2258	13253 F AT	2307	15208 S AT	2356			
2259	13344_S_AT	2308	15324 AT	2357			
2260	13370 AT	2309	15371 AT	2358			
2261	13387 AT	2310	15424 AT	2359	17566_AT		
2262	13408 S AT	2311	15463 AT	2360	17595_S_AT		
2263	13429 AT	2312	15465 AT	2361	. 17744_S_AT		
2264	13472_AT	2313	15497_S_AT	2362			
2265	13526_AT	2314	15589_S_AT	2363			
2266	13569_AT	2315	15636_S_AT	2364			
2267	13614_AT	2316	15663_S_AT	2365			
2268	13686_S_AT	2317	15770_AT	2366			
2269	13718_AT	2318	15792_AT	2367			
2270	13719_AT	2319	15855_AT	2368			
2271	13902_AT	2320	15860_AT 15891_AT	2369 2370			
2272 2273	13918_AT 13944_AT	2321 2322	15891_A1 15898 AT	2371			
2274	13944_A1 13964 AT	2322	15898_A1 15909 AT	2372			
22/4	13904_A1	2323	13303_A1	2312	. 10002_A1		

TABLE 7 (cont) 20648_S_AT 20668_AT

2426

2427

2373	18121 S AT
2374	18240 S AT
2375	18248 S AT
2376	18264_AT
2377	18276_AT
2378	18287_AT
2379	18310_AT
2380	18367_S_AT
2381	18506_AT
2382	18605_S_AT
2383 2384	18618_S_AT 18626_AT
2385	18666 S AT
2386	18834 AT
2387	18847 AT
2388	18896 AT
2389	18899 S AT
2390	18973 AT
2391	18983 S AT
2392	18988 AT
2393	18998 S AT
2394	19065_AT
2395	19119_I_AT
	19121_AT
2396	19207_AT
2397	19220_AT
2398	19284_AT
2399	19315_AT
2400	19348_AT 19403 S AT
2401 2402	19403_S_AT 19437 S AT
2402	19502 AT
2404	19609_AT
2405	19645 AT
2406	19742 AT
2407	19863 AT
2408	19873 AT
2409	19891_AT
2410	20004_S_AT
2411	20053_AT
2412	20138_AT
2413	20193_AT
2414	20199_AT
2415 2416	20220_AT 20239 G AT
2417	20297 AT
2417	20324 S AT
2419	20353 AT
2420	20362 AT
2421	20389 AT
2422	20546_AT
2423	20600_AT
2424	20623_AT
2425	20629_AT

177 TABLE 8: 2X UP IN SALT, ONLY

	LADEL	o. La or neg	LI, ONLI
12037_at	14570_at	16190_at	18506_at
12137_at	14578_s_at	16196_at	18605_s_at
12153_at	14596_at	16273_at	18626_at
12186_at	14646_s_at	16314_at	18666_s_at
12216_at	14662_f_at	16413_s_at	18747_f_at
12268_at	14668_s_at	164 1 4_at	18782_at
12449_s_at	14729_s_at	16417_s_at	18834_at
12470_at	14874_at	16455_at	18847_at
12476_at	14888_at	16548_s_at	18913_s_at
12487_at	14918_at	16582_s_at	18973_at
12493_g_at	14952_at	16589_s_at	18988_at
12609_at	14959_at	16594_s_at	18998_s_at
12685_at	14986_at	16613_s_at	19065_at
12704_f_at	15006_at	16651_s_at	19068_i_at
12709_f_at	15042_at	16668_at	19123_at
12734_f_at	15047_at	16690_g_at	19177_at
12739_s_at	15062_at	16762_at	19220_at
12750_s_at	15063_at	16820_at	19284_at
12761_s_at	15108_s_at	16873_i_at	19288_at
12819_at	15133_s_at	16987_s_at	19315_at
12845_s_at	15147_s_at	16989_at	19437_s_at
12946_at	15170_s_at	16995_at	19484_s_at
13142_at	15175_s_at	17039_s_at	19502_at
13198_i_at	15182_s_at	17040_s_at	19503_at
13229_s_at	15190_s_at	17400_s_at	19592_at
13275_f_at	15192_s_at	17425_s_at	19645_at
13344_s_at	15324_at	17433_at	19742_at
13370_at	15392_at	17467_at	19835_at
13408_s_at	15424_at	17490_s_at	19873_at
13464_at	15467_at	17529_s_at	19891_at
13472_at	15497_s_at	17543_s_at	19992_at
13526_at	15581_s_at	17566_at	20004_s_at
13614_at	15623_f_at	17595_s_at	20053_at
13652_at	15636_s_at	17744_s_at	20133_i_at
13679_s_at	15646_s_at	17758_at	20138_at
13751_at	15670_s_at	17855_at	20190_at
13918_at	15770_at	17864_at	20199_at
13919_at	15775_at	17876_at	20200_at
13944_at	15778_at	18008_r_at	20297_at
13964_at	15792_at	18013_r_at	20324_s_at
13987_s_at	15855_at	18024_s_at	20335_s_at
13993_s_at	15891_at	18027_at	20353_at
14000_at	15909_at	18053_s_at	20362_at
14032_at	15923_at	18078_at	20385_s_at
14043_at	15969_s_at	18082_at	20389_at
14052_at	15975_s_at	18090_s_at	20402_s_at
14067_at	15995_s_at	18091_at	20450_at
14070_at	15998_s_at	18121_s_at	20468_at
14269_at	16017_at	18264_at	20489_at
14285_at	1605 0_ at	18276_at	20546_at
14427_at	16067_s_at	18300_at	20569_s_at
14501_at	16072_s_at	18367_s_at	20600_at
14540_at	16165_s_at	18471_at	20623_at

20648_s_at 20678_at 20686_at 20707_s_at

178 TABLE 9: 2X UP SALT, 3 HR ONLY

20004_s_at 20053_at 20133_i_at 20138_at 20190_at

20199_at 20200_at 20220_at

20362_at 20385_s_at 20389_at 20489_at 20546_at 20623_at

20648_s_at 20678_at

20707_s_at

	TABL	
12037_at	15042_at	16987_s_at
12137_at	15047_at	16989_at
12153_at	15062_at	17039_s_at
12186_at	15063_at	17040_s_at
12216_at	15108_s_at	17425_s_at
12268 at	15133 s at	17433_at
12470 at	15147_s_at	17490_s_at
12476_at	15170_s_at	17543_s_at
12487_at	15175_s_at	17744 s at
12493 g at	15182 s at	17864 at
12609 at	15190 s at	17876 at
12685 at	15192 s at	18008 r at
12704 f at	15324_at	18013_r_at
12709 f_at	15424_at	18024_s_at
12734 f_at	15467_at	18027 at
12739 s at	15497_s_at	18053 s at
12759_s_at	15623_f_at	18078_at
12819 at	15636_s_at	18082 at
12946_at	15646 s at	18090_s_at
13142 at	15670_s_at	18091 at
		18121 s at
13229_s_at	15770_at	
13275_f_at	15775_at	18264_at
13370_at	15778_at	18276_at
13408_s_at	15792_at	18367_s_at
13464_at	15855_at	18471_at
13472_at	15891_at	18506_at
13614_at	15909_at	18605_s_at
13652_at	15923_at	18626_at
13679_s_at	15969_s_at	18666_s_at
13918_at	15975_s_at	18747_f_at
13919_at	15995_s_at	18782_at
13944_at	15998_s_at	18834_at
13987_s_at	16017_at	18847_at
13993_s_at	16050_at	18913_s_at
14000_at	16067_s_at	18973_at
14032_at	16072_s_at	18988_at
14043_at	16165_s_at	19065_at
14052_at	16196_at	19068_i_at
14067_at	16273_at	19123_at
14269_at	16314_at	19177_at
14285_at	16414_at	19220_at
14501_at	16417_s_at	19288_at
14540_at	16455_at	19315_at
14570_at	16548_s_at	19437_s_at
14596 at	16582 s at	19484_s_at
14668_s_at	16589_s_at	19502_at
14729 s at	16594 s at	19503 at
14888_at	16613 s at	19592_at
14918_at	16651_s_at	19645_at
14952 at	16668 at	19742 at
14959_at	16762 at	19835_at
14986_at	16820 at	19873 at
15006 at	16873 i at	19891 at

TABLE 10: 2X DOWN SALT, ONLY

20239 g at

20433 at

20629_at 20668_at

```
16046 s at
12011 s at
12180_at
             16060_s_at
12265 at
             16088 f at
12335 at
             16150 s at
12479 at
             16166 s at
12562 at
             16316 at
12656 at
             16340 at
12813 at
             16367_i_at
13003 s at
             16426 at
13052_s_at
             16427 at
13094_at
             16436 at
13178 at
             16489 at
             16502 at
13253 f at
13387_at
             16568_s_at
13429 at
             16638 s at
             16646_s_at
13472_at
             17273 at
13569 at
13686 s at
             17278 at
13718_at
             17567 at
13719 at
             17868 at
13902_at
             17880_s_at
14003 at
             17894 at
14144 at
             17901 at
14267_at
             17942_s_at
14418 at
             17960 at
14544 at
             17999 at
14546_s_at
             18062_at
14636 s at
             18240 s at
14951 at
             18248 s at
14956 s at
             18267 at
14979_at
             18279_s_at
14990_at
             18287_at
15040_g_at
             18310 at
15049 at
             18351_s_at
15115 f at
             18455 at
15137 s at
             18560 at
15148_s_at
             18571_at
15176 s at
             18618 s at
15208_s_at
             18896 at
15371_at
             18899_s_at
15453 s at
             18967_s_at
15463 at
             18983 s at
15465 at
             19119_i_at
15589_s_at
             19121_at
15663 s at
             19207 at
15860 at
             19348 at
15898 at
             19403 s at
15931 at
             19609 at
15965 at
             19742_at
15970 s at
             19826 at
15972_s_at
             19863 at
16005 s at
             19883 at
16028 at
             20193 at
```

TABLE 11
OSMOTIC STRESS RESPONSIVE SEQUENCES

SEQ A	FFYMETRIX	SEQ A	FFYMETRIX	SEQ Al	FFYMETRIX
ID NO:	ID NO:	ID NO:	ID NO:	ID NO:	ID NO:
2428	11994_AT	2475	13995_AT	2523	17037_S AT
2429	12028_AT	2476	14062_AT	2524	17054 S AT
2430	12033 AT	2477	14118 I AT	2525	17257_S_AT
2431	12039 AT	2478	14141 AT		18725 S AT
2432	12068 AT	2479	14310 AT	2526	17270 AT
2433	12096 AT	2480	14354 AT		17275 I AT
2434	12110 AT	2481	14476 AT	2528	17376 AT
2435	12114 AT	2482	14513_S_AT	2529	17378 AT
2436	12135_AT	2483	14568 S AT	2530	17468 AT
2437	12139 AT	2484	14604 AT	2531	17481 AT
2438	12189 AT	2485	14634 S AT	2532	17511_S_AT
2439	12191 AT	2486	14660 S AT	2533	17519 S AT
2440	12211 AT	2487	14666 S AT	2534	17815 S AT
2441	12223 S AT	2488	14686 S AT	2535	17897 AT
2442	12366 S AT		17464_AT	2536	17923 S AT
	12869 S AT	2489	14726 S AT	2537	17934 AT
2443	12381 AT	2490	14848 S AT	2538	17937 S AT
2444	12406 S AT	2491	14873 AT	2539	17944 AT
2445	12412 AT	2492	14883 AT	2540	17958 AT
2446	12453 AT	2493	15082 AT	2541	18216 AT
2447	12571 S AT	2494	15121 S_AT	2542	18227 AT
2448	12662 AT		16014 S AT	2543	18284 AT
2449	12746 I AT	2495	15168 S AT	2544	18301 S AT
2450	12774 AT	2496	15271 AT	2545	18312 S_AT
2451	12787 AT	2497	15338 AT	2546	18326 S AT
2452	12847 AT	2498	15418 AT	2547	18369 AT
2453	12848 AT	2499	15429 AT	2548	18411 AT
2454	12895_AT	2500	15548 AT	2549	18533 AT
2455	12911 S AT	2501	15666 S AT	2550	18576 S AT
2456	12920 AT	2502	15672 S AT	2551	18599 AT
	12921 S AT	2503	15680_S_AT	2552	18640_AT
2457	13027_AT	2504	15867 AT	2553	18672 S AT
2458	13059 AT	2505	15918 AT	2554	18720 S AT
2459	13075 I AT	2506	15999 S AT	2555	18768_AT
2460	13180 S AT	2507	16303 AT	2556	18877_AT
2461	13255 I AT	2508	16363 AT	2557	18942_AT
2462	13270 AT	2509	16440 S AT	2558	18945_AT
	18167 S AT	2510	16458 S AT	2559	18960 AT
2463	13283 S AT	2511	16475 AT	2560	18965 AT
2464	13382 AT	2512	16513 S AT	2561	19060 AT
2465	13386 S AT	2513	16529_AT	2562	19164_G_AT
2466	13433 AT	2514	16547 S AT	2563	19266 AT
2467	13482 AT	2515	16553 F AT	2564	19366 S AT
2468	13732 AT	2516	16563 S AT	2565	19369_AT
2469	13733_I_AT	2517	16629_S_AT	2566	19371_AT
2470	13842_AT	2518	16797_AT	2567	19386_AT
2471	13860_S_AT	2519	16814_AT	2568	19412_AT
2472	13868_AT	2520	16832_AT	2569	19427_S_AT
2473	13901_AT	2521	16976_S_AT	2570	19622_G_AT
2474	13933_AT	2522	17007_AT	2571	19681_AT

TABLE 11 (cont)

2572	19819_S_AT
2573	19961_S_AT
2574	20002_AT
2575	20034 I AT
2576	20062_AT
2577	20136_AT
	20223_AT
2579	20235_I_AT
2580	20401_AT
	20407_AT
2582	20470_AT
2583	20626_AT
2584	20631_S_AT
2585	20647_AT

TABLE 12: 2X UP IN MANNITOL, ONLY

12039 at 16832 at 12068_at 16993_at 12139 at 17037 s at 12212_at 17054_s_at 12278_at 17083 s at 12366_s_at 17097_s_at 12453 at 17119 s at 12556 at 17270 at 17305 at 12575 s at 12746 i at 17376 at 17378 at 12848 at 12869 s at 17449 s at 12920 at 17481 at 12921 s at 17533 s at 13041 s at 17832 s at 13059 at 17923 s at 13241_s_at 17944 at 13255 i at 18059 i at 13270_at 18216 at 13382 at 18230 at 13406 at 18255 at 13433 at 18284 at 18301_s_at 13550 at 13672 s at 18312 s at 13716_at 18326 s at 13842 at 18599 at 13933 at 18672_s_at 13995 at 18720 s at 14062 at 18768 at 14075 at 18814 at 18877 at 14162 at 14208 at 18921_g_at 14217 at 18960 at 14235 at 19060 at 14310_at 19182_at 14431 at 19192 at 19266_at 14513_s_at 14584 at 19369 at 14604 at 19386 at 14673 s at 19402_at 14856 s at 19412 at 15207_s_at 19432 s at 15338 at 19469 at 15406 at 19622 g at 15418_at 19819 s at 15591_s_at 19826 at 15666_s_at 20152_at 15680_s_at 20223 at 15866 s at 20235 i at 15918_at 20365 s at 16340 at 20470 at 16553 f at 20537 at

16797 at

20547 at

TABLE 13: 2X UP IN MANNITOL, 3 HR ONLY

	IAC
12039_at	17449_s_at
12068_at	17481_at
12139_at	17533_s_at
12212_at	17923_s_at
12278_at	17944_at
12366_s_at	18059_i_at
12453_at	18216_at
12556_at	18230_at
12575_s_at	18255_at
12746_i_at	18301_s_at
12848_at	18312_s_at
12869_s_at	18326_s_at
12920_at	18599_at
12921_s_at	18720_s_at
13041_s_at	18768_at
13059_at	18814_at
13241_s_at	18877_at
13382_at	18921_g_at
13406_at	18960_at
13433_at	19060_at
13550_at	19192_at
13672_s_at	19266_at
13933_at	19369_at
13995_at	19386_at
14062_at	19402_at
14075_at	19412_at
14162_at	19432_s_at
14217_at	19469_at
14310_at	19622_g_at
14431_at	19819_s_at
14513_s_at	20152_at
14584_at	20223_at
14604_at	20235_i_at
14673_s_at	20365_s_at
14856_s_at	20470_at
15207_s_at	20537_at
15338_at	
15418_at	
15591_s_at	
15866_s_at	
15918_at	
16340_at	

16553 f_at 16797_at 16832_at 17037_s_at 17054_s_at 17083_s_at 17097_s_at 17270_at 17305_at 17376_at 17378_at

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TABLE 14: 2X DOWN IN MANNITOL, ONLY

	TABLE 14
12028 at	14897 at
12033_at	14918 at
12110_at	15082 at
12114 at	15084 at
12189 at	15098 s at
12191 at	15105 s at
12211 at	15121 s at
12223 s at	15126_s_at
12268 at	15168_s_at
12345 at	15271 at
12381 at	15429 at
12406 s at	15548 at
12412 at	15672 s_at
12522 at	15753 at
12571_s_at	15867 at
12662 at	15999_s_at
12787 at	16001 at
12767_at	
12895_at	16190_at
12911_s_at	16260_at
13027_at	16303_at
13075_i_at	16363_at
13221_at	16458_s_at
13262_s_at	16468_at
13283_s_at	16475_at
13386_s_at	16513_s_at
13447_s_at	16529_at
13482_at	16563_s_at
13634_s_at	16690_g_at
13709_s_at	16814_at
13732_at	16847_at
13733_i_at	16927_s_at
13812_s_at	16976_s_at
13825_s_at	17007_at
13860_s_at	17014_s_at
13868_at	17016_s_at
13901_at	17071_s_at
14052_at	17090_s_at
14224_at	17257_s_at
14244_s_at	17275_i_at
14254_s_at	17424_at
14256_f_at	17464_at
14354_at	17468_at
14476_at	17511_s_at
14568_s_at	17519_s_at
14634_s_at	17525_s_at
14646_s_at	17645_s_at
14660_s_at	17741_at
14686_s_at	17815_s_at
14726_s_at	17897_at
14848_s_at	17899_at
14873_at	17934_at
14883_at	17937_s_at
	· ·

17958 at 18012_s_at 18227 at 18272 at 18331 s at 18369 at 18411 at 18533_at 18576 s at 18640 at 18696 s at 18945 at 18949_at 18953 at 18965_at 19164 g at 19322 at 19366 s at 19371_at 19397_at 19427_s_at 19681_at 19707 s at 19839 at 19961_s_at 19976 at 19998 at 20002 at 20034 i at 20136_at 20382 s at 20407 at 20529_at 20626 at 20631_s_at 20647_at 20699 at

TABLE 15
COLD & OSOMOTIC STRESS RESPONSIVE SEQUENCES

				_	
SEQ A	FFYMETRIX	SEO A	FFYMETRIX	SEO A	FFYMETRIX
ID NO:	ID NO:	ID NO:	ID NO:	ID NO:	ID NO:
1699	12040 AT	1742	13262 S AT	1787	14431 AT
1700	12048 AT	1743	13286 S AT	1788	14480 AT
1701	12054 S AT	1744	13324 AT	1789	14497 AT
1702	12077 AT	1745	13340 S AT	1790	14553 AT
1703	12107 I AT	1746	13361 AT	1791	
1703	12107_1_A1 12113_AT	1747	13406 AT	1791	14584_AT
1704	12115_A1 12154 AT	1747			14600_AT
1706		1749	13441_S_AT	1793	14673_S_AT
	12171_AT		13513_AT	1504	19432_S_AT
1707	12212_AT	1750	13550_AT	1794	14681_G_AT
1708	12278_AT	1751	13573_AT	1795	14699_AT
1709	12317_AT	1752	13577_S_AT	1796	14751_AT
1710	12325_AT	1753	13606_AT	1797	14762_AT
1711	12333_AT	1754	13609_AT	1798	14828_S_AT
1712	12345_AT	1755	13625_S_AT	1799	14856_S_AT
1713	12349_S_AT	1756	13626_AT	1800	14882_AT
	14254_S_AT	1757	13634_S_AT	1801	14897_AT
	14256_F_AT	1758	13672_S_AT	1802	14978_AT
1714	12356 AT		18916 S AT	1803	14985 S AT
1715	12380 AT	1759	13709 S AT	1804	15031 AT
1716	12392 AT	1760	13736 AT	1805	15084 AT
1717	12460 S AT	1761	13775 AT	1806	15096 AT
1718	12556 AT	1762	13810 AT	1807	15105 S AT
1719	12575 S AT	1763	13812 S AT	1808	15110 S AT
1720	12686 S AT	1764	13825 S AT	1809	15111 S AT
1721	12701 I AT	1765	14015 S AT	1810	15120_S_AT
1722	12754 G AT		14016 S AT	1811	15126 S AT
1723	12782 R AT	1766	14029 AT	1812	15142 S AT
1724	12784 AT	1767	14036 AT	1813	15144 S AT
1725	12879 S AT	1768	14051 AT	1814	15184 S AT
1726	12891 AT	1769	14060 AT	1815	15194_S_AT
1720	16817 S AT	1770	14064 AT	1816	15203 S AT
1707	12898 G AT	1770	14064_AT	1817	15203_S_AT 15207 S AT
1727					
1728	12974_AT	1772	14075_AT	1818	15240_AT
1729	12998_AT	1773	14094_S_AT	1819	15366_AT
1730	13041_S_AT	1004	19999_S_AT	1820	15398_AT
1731	13124_AT	1774	14096_AT	1821	15406_AT
1732	13134_S_AT	1775	14104_AT	1822	15448_AT
1733	13144_AT	1776	14123_S_AT	1823	15466_AT
1734	13147_AT	1777	14126_S_AT	1824	15481_AT
1735	13152_S_AT	1778	14131_AT	1825	15484_AT
1736	13187_I_AT	1779	14136_AT	1826	15549_AT
	16981_S_AT	1780	14139_AT	1827	15591_S_AT
1737	13192_S_AT		14140_AT	1828	15606_S_AT
	17525_S_AT	1781	14162_AT	1829	15614_S_AT
1738	13212_S_AT		14217_AT		16927_S_AT
		1782	14178_AT	1830	15629_S_AT
1739	13215_S_AT	1783	14201_AT	1831	15633_S_AT
	16649_S_AT	1784	14208_AT	1832	15641_S_AT
1740	13241_S_AT	1785	14235_AT		18012 S AT
1741	13246_AT	1786	14242_S_AT	1833	15720_AT
	_				_

TABLE 15 (cont)

1834	15815 S AT	1884	17452_G_AT	1936	19469 AT
1835	15817 AT	1885	17540 S AT	1937	19409_AT
1836	15837 AT	1886	17552 S AT	1938	19597_S AT
1837	15841 AT	1887	17571 AT	1939	19710 S AT
1838	15866 S AT	1888	17589 AT	1940	19830 AT
1000	18255 AT	1889	17641 G AT	1940	19839 AT
1839	15872 AT	1890	17741 AT	1941	19840 S AT
1039	18331 S AT	1070	18098 AT	1942	19840_S_AT
1840	15892_AT	1891	17766_AT	1944	19860_AT
1841	15933 AT	1892	17873 S AT	1944	19880 AT
1842	15947 AT	1893	17904 AT	1946	19889 AT
1843	15959 S AT	1894	17920 S AT	1947	19898 AT
1844	16001 AT	1895	17920_3_AT 17925_AT	1948	19914 AT
1845	16052 AT	1896	17943 AT	1949	19914_AT
1846	16161 S AT	1897	18059 I AT	1950	19924_AT 19949_AT
1847	16204_AT	1898	18230 AT	1951	19949_AT
		1899		1951	19976_AT 19998 AT
1848	16232_S_AT		18263_AT		
1849	16252_AT	1900	18272_AT	1953	20030_AT
1850	16260_AT	1901	18540_AT	1954	20151_AT
1851	16266_AT	1902	18608_AT	1955	20152_AT
1852	16299_AT	1903	18647_AT	1956 1957	20187_AT
1853	16365_AT	1904	18662_S_AT		20214_I_AT
1854	16468_AT	1905	18664_AT	1958	20269_AT
1855	16477_AT 16491 AT	1906	18695_S_AT 18704_AT	1959 1960	20271_AT 20273_AT
1856		1907		1961	
1857	16523_S_AT	1908 1909	18814_AT 18907 S AT	1961	20299_AT 20323_AT
1858	16566_S_AT	1910		1962	20429_S_AT
1859 1860	16570_S_AT 16688 AT	1910	18921_G_AT 18924_AT	1964	20429_S_AT
1861	16840 AT	1911	18949 AT	1965	20480 S AT
1862	16847 AT	1912	19707 S AT	1966	20529_AT
1863	16893 AT	1913	18995 AT	1967	20547 AT
1864	16896 S AT	1914	19017 AT	1968	20555 S AT
1865	16898 S AT	1915	19034 AT	1969	20699_AT
1866	16912 S AT	1916	19063 AT	1707	20077_11
1867	16980 AT	1917	19142 AT		
1868	16993 AT	1918	19158 AT		
1869	17008 AT	1919	19180 AT		
1870	17012 S AT	1920	19187 AT		
1871	17012_S_AT	1921	19192 AT		
1872	17014_S_AT	1922	19195 AT		
1873	17032 S AT	1923	19199 AT		
1874	17050_S_AT	1924	19231 AT		
1074	17051 S AT	1925	19263 AT		
1875	17071 S AT	1926	19308 AT		
1876	17090 S AT	1927	19322 AT		
10.0	18690 S AT	1928	19365 S AT		
1877	17097 S AT	1929	19372 AT		
1878	17104 S AT	1930	19389_AT		
1879	17119 S AT	1931	19392 AT		
1880	17160 AT	1932	19397 AT		
1881	17305 AT	1933	19400_AT		
1882	17424 AT	1934	19402 AT		
1883	17449 S AT	1935	19458 AT		
_000					

TABLE 16: 2X UP IN MANNITOL & COLD, ONLY

12345 at	
12784 at	
13153_r_at	
13212_s_at	
13215_s_at	
13246 at	
13262_s_at	
13361 at	
13625 s at	
13764 at	
13810 at	
14015_s_at	
14016_s_at	
14060 at	
14096 at	
14123 s at	
14139_at	
14219_at	
14248 at	
14254_s_at	
14256_f_at	
14609 at	
14636_s_at	
14681_g_at	
14699 at	
14704_s_at	
14828 s at	
14882_at	
15110 s at	
15184 s at	
15448_at	
15629 s at	
15720 at	
15846 at	
15947_at	
16161_s_at	
16365 at	
16427 at	
16566_s_at	
16570 s at	
16649_s_at	
16688_at	
16712 at	
16817_s_at	
16840_at	
16893_at	
16912_s_at	
16916_s_at	
16927_s_at	
16981_s_at	
17012_s_at	
17014_s_at	
17051 c ot	

17051_s_at

17066 s at 17540 s at 17567_at 17766 at 17904_at 17920 s at 17943 at 18263 at 18351_s_at 18662 s at 18670 g_at 18695_s_at 18704_at 18729 at 18995 at 19158 at 19473_at 19710_s_at 19883_at 19889 at 20030_at 20269 at 20271_at 20299_at 20429_s_at 20438 at 20480_s_at

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	TABLE 17: 2X	DOWN COL
12040 at	14553 at	17873 s at
12048_at	14612_at	17925_at
12054_s_at	14751 at	18098 at
12077_at	14762 at	18540 at
12107 i at	14978_at	18608_at
12113 at	14985 s at	18647_at
12154 at	15031 at	18664 at
12171 at	15096 at	18690_s_at
12317 at	15111 s at	18725 s at
12325_at	15120_s_at	18924_at
12333 at	15142 s at	19017 at
12356 at	15198 s at	19034 at
12380_at	15203_s_at	19063_at
12392_at	15240_at	19141_at
12460 s at	15366 at	19142 at
12686 s at	15392 at	19180_at
12701_i_at	15398_at	19187_at
12782 r at	15466_at	19195 at
12879_s at	15481 at	19199 at
12898 g at	15484 at	19231 at
12974_at	15549_at	19308_at
12998 at	15623_f_at	19372_at
13144_at	15815_s_at	19392 at
13147_at	15817_at	19392_at
13152 s at	15841 at	19458_at
13192_s_at	15892 at	19597 s at
13286_s_at	15933_at	19762 at
13324 at	15959_s_at	19830 at
13340 s at	16052 at	19853_at
13441_s_at	16204_at	19869 at
13513_at	16252_at	19880_at
13573 at	16266_at	19898 at
13606_at	16299 at	19914_at
13609_at	16477_at	19924_at
13626_at	16491 at	19949 at
13736 at	16561 s at	20151 at
13775_at	16645_s_at	20131_at
14029_at	16898 s at	20214 i at
14036_at	16980_at	20273 at
14051_at	17008 at	20323 at
14064_at	17104_s_at	20457_at
14066 at	17160_at	20555_s_at
14094 s at	17317 at	20000_0_0
14104_at	17400_s_at	
14126 s at	17452_g_at	
14131_at	17477 s at	
14136 at	17500_s_at	
14178_at	17552 s at	
14192 at	17571 at	
14201_at	17572 s at	
14242 s_at	17589 at	
14480 at	17641_g_at	
14497_at	17855_at	
at	005_01	

TABLE 18

COLD & SALINE STRESS RESPONSIVE SEQUENCES

oro i	DENA CETDIN	2010	10544 AT	2062	15047 AT
	FFYMETRIX	2018	13544_AT	2062	15047_AT
ID NO:	ID NO:	2019	13549_AT	2063	15063_AT
1970	12021_AT	2020	13565_AT	2064	15085_S_AT
1971	12037_AT		FFYMETRIX	2065	15123_S_AT
1972	12094_AT	ID NO:	ID NO:	2066	15133_S_AT
1973	12098_AT	2021	13580_AT	2067	15137_S_AT
1974	12128 AT	2022	13588 AT	SEQ A	FFYMETRIX
1975	12148 AT	2023	13649 AT	ID NO:	ID NO:
1976	12151 AT	2024	13652 AT	2068	15153 S AT
1977	12357 S AT	2025	13679 S AT	2069	15170 S AT
1978	12394 AT	2026	13696 AT	2070	15172 S AT
1979	12472 S AT	2027	13702 S AT	2071	15182 S AT
1980	12475 AT	2028	13751 AT	2072	15190 S AT
1981	12482 S AT	2029	13919 AT	2073	15241 S AT
1982	12490_AT	2030	13943 AT	2074	15389 AT
1983	12505 S AT	2031	13950 S AT	2075	15453 S AT
		2032	14050 AT	2076	15495 AT
1984	12531_AT	2032		2077	15496 AT
1985	12540_S_AT	2033	14055_S_AT		
1986	12541_AT		16166_S_AT	2078	15519_S_AT
1987	12577_AT	2034	14067_AT	2079	15562_AT
1988	12594_AT	2035	14078_AT	2080	15580_S_AT
1989	12629_AT	2036	14110_I_AT	2081	15582_S_AT
1990	12642_AT	2037	14144_AT	2082	15638_S_AT
1991	12656_AT	2038	14232_AT		18751_F_AT
1992	12660 AT	2039	14285_AT	2083	15646_S_AT
1993	12712 F AT	2040	14346_AT	2084	15647_S_AT
1994	12725 R AT	2041	14432_AT	2085	15654_S_AT
1995	12745 AT	2042	14468 AT	2086	15655_S_AT
1996	12777 I AT	2043	14479 AT	2087	15658 S_AT
1997	12790_S_AT	2044	14524 S AT	2088	15670_S_AT
1998	12798 AT	2045	14608 AT	2089	15775 AT
1999	12801 AT	2046	14621 AT	2090	15798 AT
2000	12855 F AT	2047	14635 S AT	2091	15930 AT
2001	12887 S AT		17128 S AT	2092	15931 AT
2002	12933 R AT	2048	14640 S AT	2093	15949 S AT
2003	12951 AT	2049	14643 S AT	2094	16017 AT
2004	13005 AT	2050	14663_S_AT	2095	16053 I AT
2005	13015 S AT	2051	14668 S AT	2096	16078 S AT
2006	13115 AT	2052	14688 S AT	2097	16086 S AT
2007	13178 AT	2002	18279_S_AT	2098	16120_S_AT
2007	13228 AT	2053	14737 S AT	2099	16126 S AT
2009		2054	14768 AT	2100	16150 S AT
2009	13236_S_AT	2055	14875 AT	2101	16159 S AT
-040	16646_S_AT	2056	14911 S AT	2102	16230 AT
2010	13266_S_AT	2030		2102	16306 AT
	15211_S_AT	2057	17056_S_AT		
2011	13275_F_AT	2057	14924_AT	2104	16367_I_AT
2012	13335_AT	2058	14956_S_AT	2105	16417_S_AT
2013	13362_S_AT		15148_S_AT	2106	18083_R_AT
2014	13428_AT		18673_AT	2106	16418_S_AT
2015	13464_AT	2059	14964_AT	2107	16423_AT
2016	13480_AT	2060	15022_AT	2108	16449_S_AT
2017	13538_AT	2061	15040_G_AT	2109	16484_S_AT

TABLE 18 (cont)

		***	522 IS (1011)
2110	16489 AT	2163	18455 AT
2111	16565 S AT	2164	18459 AT
2112	16596 S AT	2165	18571 AT
2113	16600 S AT	2166	18604 AT
2114	16603 S AT	2100	19181 S AT
		2167	18644 AT
2115	16638_S_AT		
2116	16642_S_AT	2168	18745_F_AT
2117	16763_AT		19611_S_AT
2118	16914_S_AT	2169	18782_AT
2119	16968_AT	2170	18881_AT
2120	16983_AT	2171	18904_S_AT
2121	16989_AT	2172	18914_S_AT
2122	17002_AT	2173	18963_AT
2123	17015_S_AT	2174	19068_I_AT
2124	17040_S_AT	2175	19078_AT
	18913_S_AT	2176	19171_AT
2125	17232_AT	2177	19177_AT
2126	17380_AT	2178	19394_AT
2127	17394_S_AT	2179	19411_AT
	20640_S_AT	2180	19415_AT
2128	17398_AT	2181	19466_S_AT
2129	17448_AT	2182	19484_S_AT
2130	17485_S_AT	2183	19549_S_AT
2131	17490_S_AT	2184	19592_AT
2132	17499_S_AT	2185	19633_AT
2133	17505_S_AT	2186	19641_AT
2134	17516_S_AT	2187	19669_AT
2135	17529_S_AT	2188	19672_AT
2136	17543_S_AT	2189	19684_AT
2137	17593_R_AT	2190	19692_AT
	19858_S_AT	2191	19746_AT
2138	17609_AT	2192	19835_AT
2139	17698_AT	2193	19848_S_AT
2140	17836_AT	2194	19892_AT
2141	17886_AT	2195	19904_AT
2142	17896_AT	2196	19936_AT
2143	17901_AT	2197	19974_S_AT
2144	17902_S_AT	2198	19994_AT
2145	17913_S_AT	2199	20005_S_AT
2146	17924_AT	2200 2201	20022_AT 20032_AT
2147	17954_S_AT 17960 AT	2201	20032_AT
2148		2202	20044_AT
2149	17991_G_AT 18967 S AT	2203	20049_A1 20081 AT
2150		2204	20133 I AT
2150	17999_AT	2206	20155 S AT
2151 2152	18057_I_AT 18078_AT	2207	20163 S AT
2152	18091 AT	2208	20200 AT
2154	18168 S AT	2209	20296 S AT
2154	18252 AT	2210	20336 AT
2156	18267 AT	2211	20341_AT
2157	18300 AT	2212	20372 AT
2158	18308 I AT	2212	20385 S AT
2159	18328 AT	2214	20433 AT
2160	18354 AT	2215	20489 AT
2161	18402 AT	2216	20525 AT
2162	18416 AT	2217	20543 AT
2102		2217	

191 TABLE 19: 2X UP IN SALT & COLD, ONLY

TABLE 19:	2X UP IN SA
15495_at	18745 f at
15496_at	18904 s at
15519_s_at	18914_s_at
15580 s at	18929 s at
15582 s_at	18946_at
15776 at	18963 at
15798 at	19078 at
15910 at	19137_at
	19141 at
15937 at	19411 at
15949 s at	19641 at
15972 s at	19672 at
16048 at	19684_at
16086 s at	19692 at
16120 s at	19746 at
	19762 at
	19869_at
16159 s at	19894 at
	19904 at
	19936 at
	19994 at
	20005 s_at
	20031_at
16565 s at	20044 at
16603 s at	20382 s at
	20406 q at
16968_at	20421_at
16983_at	20525_at
17002_at	20543_at
17015_s_at	20565_at
17019_s_at	20570_at
	20640_s_at
17232_at	20646_at
17317_at	20720_at
10484_at	
	15495_at 15496_at 15496_at 15519_s_at 15580_s_at 15580_s_at 15580_s_at 15580_s_at 15776_at 15791_at 15931_at 15937_at 15937_at 15949_s_at 15949_s_at 16048_at 16086_s_at 16150_s_at 16150_s_at 16150_s_at 16443_at 16449_s_at 16449_s_at 166565_s_at 166565_s_at 166565_s_at 166663_s_at 166663_s_at 166663_s_at 166663_s_at 16763_at 16968_at 17002_at 17015_s_at 17015_s_at 17015_s_at 170179_s_at 17078_s_at 1708_s_at 1708_s_a

19394 at 19415_at 19466 s at 19549_s_at 19592 at 19633_at 19669 at 19848_s_at 19858 s at 19878_at 19892_at 19974 s at 20022_at 20032_at 20049 at 20081 at 20155_s_at

20163_s_at 20296_s_at 20336_at 20341_at 20365_s_at 20372_at 20489_at 20491_at 20577_at 20609_at 20672_at

TABLE 20: 2X DOWN IN COLD & SALT, ONLY

	TABLE 20:
12021 at	15123_s_at
12094 at	15153 s at
12128 at	15172 s at
12151 at	15190 s at
12332 s_at	15211 s at
12472 s at	15241 s at
12475 at	15437 at
12482 s at	15562 at
12490 at	15638 s at
12531 at	15647 s at
12540 s at	15654 s at
12577 at	15655 s at
12629 at	15658 s at
12642 at	15695 s at
12660 at	15846 at
12676 s at	15930 at
12712 f at	16053 i at
12725 r at	16078 s at
12777 i at	16229 at
12777_1_at	16465 at
12801 at	16484 s at
12887 s at	16596 s at
	16600 s at
	16642 s at
	16914 s at
13228_at 13362 s at	
	17027_s_at 17066 s at
13428_at 13538_at	17080_s_at
13565 at	17128 s at
13588 at	17380 at
13696 at	17398 at
13702 s at	17448 at
13716 at	17485 s at
13764 at	17490 s at
14050 at	17490_3_at
14055_at	17505 s at
14069 at	17503_s_at
14078 at	17593_r_at
14232 at	17886 at
14346 at	17913 s at
14608 at	17924 at
14609 at	17954 s at
14621 at	17991 g at
14635 s at	18057 i at
14663 s at	18069 at
14688 s at	18328 at
14691 at	18416 at
14704 s at	18604 at
14704_s_at	18644 at
14911 s at	18881 at
14964 at	19171 at
15022 at	19181 s at
15085 s at	19182 at
15065_5_at	13102_at

SEQ AFFYMETRIX ID NO:

2685

2686

2687

2688

2689

2690

2691

2692

2693

2694

2695

2696

2697

2698

2699

2700

2701

2702 2703

ID NO: 2681 19409 AT 2682 19503 AT 2683 19826 AT 2684

19847 S AT

19930 AT

19992 AT

20096 AT

20108_AT

20256_S_AT

20290 S AT

20298 AT

20305 AT 20322 AT

20333 AT

20424_AT

20450 AT

20468_AT

20639 AT

20678 AT

20686 AT

20402 S AT

20446_S_AT

20569_S_AT

TABLE 21 OSMOTIC & SALINE STRESS RESPONSIVE SEQUENCES

aro.	A PENA (PED IN	aro	A DESCRIPTION
	AFFYMETRIX		AFFYMETRIX
ID NO:	ID NO:	ID NO:	ID NO:
2586	12126_S_AT	2634	16073_F_AT
2587	12126_S_AT 12137_AT	2635	16114_S_AT
2588	12227_AT 12239_AT	2636	16127_S_AT
2589	12239_AT		18744_F_AT
2590	12268_AT 12369_AT 12476_AT	2637	16190_AT
2591	12369_AT	2638	16196_AT
2592	12476 AT	2639	16236 G AT
2593	12484_G_AT 12494_AT		19531_AT
2594	12494 AT	2640	16310_AT
2595	12644_AT 12645_AT	2641	16316 AT
2596	12645 AT	2642	16334 S AT
2597	12796 S AT	2643	16335 AT
2596 2597 2598	12819 AT	2644	16340 AT
2599	12841 AT	2645	16450 S AT
2600	12852 S AT	2646	16500_AT
2000	19455_S_AT	2647	16524 AT
2601		2640	16533 AT
2601 2602	13171 AT	2649	16690 G AT
2603	13174 R AT	2650	16762 AT
2604	13596 AT	2651	16819 AT
2604		2652	16873 I AT
	13807_AT	2652 2653 2654	16972 AT
2606 2607	13977_AT	2053	
		2654	16991_AT
2608 2609	14052_AT	2655 2656	17099_S_AT
		2656	17339_AT
2610	14335_AT	2657	17397_S_AT
2611	14486_AT	2658 2659	17419_AT
2612	14506_AT		
2613 2614	14518_AT	2660 2661	17554_S_AT
			17939_AT
2615	14578_S_AT	2662	18013_R_AT
2616	14646_S_AT		18178_S_AT
2617	14662_F_AT	2663	18024_S_AT
	15962_S_AT	2664	18032_I_AT
2618	14901_AT	2665	18054_AT
2619	14918 AT	2666	18151_AT
2620	14986_AT	2667	18281_AT
2621	15053 S AT	2668	18445 AT
2622	15179 S AT	2668 2669 2670	18520 AT
2623	15252 G AT	2670	18583 AT
2624	15280 AT	2671	18663 S AT
2625	15467_AT	2671 2672	18753 S AT
2626	15607 S AT	2673	18876 AT
2627	15625 S AT	2672 2673 2674 2675	18938 G AT
2628	15703 I AT	2675	18971 AT
2629	15827 AT	2676	18977 AT
2630	15863 AT	2676 2677	18981 AT
2631	15022 AT	0.000	10000 475
2632	15946_S_AT 16005_S_AT	2678	19196 AT
2633	16005 S AT	2680	19376 AT
2000	10000_0_111		

TABLE 22: 2X UP IN SALT & MANNITOL, ONLY

	TABLE 22
12126_s_at	17548_s_at
12227_at	17554_s_at
12369_at	17961_at
12521 at	18032 i at
12644_at	18054_at
12645_at	18151_at
12724_f_at	18167_s_at
12795_at	18281_at
12796_s_at	18520_at
12841_at	18663_s_at
12852_s_at	18744_f_at
12958_at	18753_s_at
13014_at	18789_at
13174_r_at	18876_at
13211_s_at	18909_s_at
13596_at	18938_g_at
13640_at	18977_at
13789_at	19099_at
13977_at	19108_at
13999_at	19135_at
14069_at	19227_at
14083_at	19376_at
14089_at	19429_at
14293_at	19455_s_at
14675_s_at	19531_at
15053_s_at	19789_s_at
15058_s_at	19878_at
15252_g_at	20017_at
15280_at	20096_at
15437_at	20256_s_at
15607_s_at	20290_s_at
15625_s_at	20305_at
15827_at	20322_at
15863_at	20333_at
15880_at	20420_at
16005_s_at	20424_at
16031_at	20689_s_at
16073_f_at	
16316_at	
40004 - 4	

16334_s_at 16335_at 16450_s_at 16500_at 16524_at 16533_at 16597_s_at 16819_at 17085_s_at 17099_s_at 17419_at 17442_i_at

TABLE 23: 2X DOWN IN MANNITOL & SALT, ONLY

12239_at 20108_at 12251_at 20298_at 12476_at 20421_at 12484_g_at 20432_at 12561_at 20639_at 12647_s_at

12561_at 12647_s_at 12719_f_at 12819_at

12841_at 13084_at 13171 at

13172_s_at 13435_at

13807_at 14250 r_at

14230_1_at

14486_at 14506_at

14518_at 14901_at

15046_s_at 15179_s_at

15451_at 15703_i_at

15946_s_at 16014 s at

16114_s_at 16310_at

16342_at 16712_at

16712_at

16972_at 16991_at

17397_s_at 17408_at

17460_at 17775 at

17939_at 18445_at

18583_at

18751_f_at 18971_at

18981_at 19156 s at

19196_at

19359_s_at 19409_at

19503_at 19713_at

19718_at 19847 s at

19930_at

TABLE 24
COLD, OSMOTIC & SALINE RESPONSIVE SEQUENCES

ero.	AFFYMETRIX	SEQ .	AFEVMETRIV	ero.	A PENA APERDINA
			AFFYMETRIX		AFFYMETRIX
ID NO:	ID NO:	ID NO:	ID NO:	ID NO:	ID NO:
1262	12004_AT	1306	12945_AT	1347	13725_AT
1263	12023_S_AT	1307	12958_AT	1348	13764_AT
1264	12078_AT	1308	12964_AT	1349	13771_AT
1265	12115_AT	1309	12968_AT	1350	13789_AT
1266	12118_AT	1310	12972_AT	1351	13916_AT
1267	12150_AT	1311	12989_S_AT	1352	13965_S_AT
1268	12251_AT	1312	13004_AT	1353	13967_AT
1269	12271_S_AT	1313	13014_AT	1354	14028_AT
1270	12276 AT	1314	13025 AT	1355	14039 AT
1271	12332 S AT	1315	13036 AT	1356	14046 AT
	13211 S AT	1316	13099 S AT	1357	14049 AT
1272	12338 AT	1317	13136 AT	1358	14069 AT
1273	12400 AT	1318	13146 S AT	1359	14077 AT
1274	12430 AT		13239 S_AT	1360	14080 AT
1275	12457 AT	1319	13153 R AT	1361	14083 AT
1276	12521 AT	1320	13159 AT	1362	14089 AT
1277	12522 AT	1321	13176 AT	1363	14090 I AT
1278	12530 AT	1322	13217 S AT	1364	14097 AT
1279	12536 S_AT	1322	17500 S AT	1365	14116 AT
1279	12538_AT	1323	13225 S AT	1366	14110_AT
1281		1323	15997 S AT	1300	14131_A1 14219 AT
	12561_AT	1324	13230 S AT	1367	14219_AT 14170 AT
1282	12574_AT	1324		1368	
	19019_I_AT	1205	15972_S_AT		14172_AT
1283	12595_AT	1325	13279_S_AT	1369	14192_AT
1284	12606_AT		17477_S_AT	1370	14224_AT
1285	12609_AT	1326	13280_S_AT	1371	14227_AT
1286	12622_AT		20301_S_AT	1372	14244_S_AT
1287	12630_AT	1327	13282_S_AT		14245_AT
1288	12647_S_AT		17027_S_AT		14645_S_AT
1289	12676_S_AT	1328	13426_AT		15974_G_AT
1290	12697_AT	1329	13432_AT	1373	14248_AT
1291	12698_AT	1330	13435_AT	1374	14250_R_AT
1292	12719_F_AT	1331	13447_S_AT	1375	14367_AT
1293	12724_F_AT	1332	13474_AT	1376	14381_AT
	15871_S_AT	1333	13511_AT	1377	14384_AT
	16597_S_AT	1334	13546_AT	1378	14398_S_AT
1294	12749 AT	1335	13547_S_AT	1379	14487_AT
1295	12765 AT	1336	13548 AT	1380	14582_AT
1296	12769 AT	1337	13555 AT	1381	14597 AT
1297	12781 AT	1338	13587 AT	1382	14609 AT
1298	12785 AT	1339	13595 AT	1383	14612 AT
1299	12792 S AT	1340	13610 S AT		19267 S AT
1300	12795 AT	1341	13627 AT	1384	14614 AT
1301	12805 S AT	1342	13640 AT	1385	14636 S AT
1302	12857 AT	1343	13645 AT	1386	14644 S AT
1303	12883 S AT	1344	13647 AT	20	14658 S AT
1304	12909 S AT	1345	13706 S AT		14659 S AT
	16539 S AT	10	19701 S AT		15964 S AT
1305	12932 S AT	1346	13716 AT	1387	14675 S AT
1505	15605 S AT	15.10	18228 AT	150,	1.5.5_5_111
	15.555_5_111				

TABLE 24 (cont)

1388	14691 AT	1443	15753 AT	1496	16789 AT
	14709 AT	1444	15761 AT	1497	16818 S AT
1389	14704 S AT	1445	15776 AT	1498	16971 S AT
	15846 AT	1446	15778 AT	1499	17018 S AT
1390	14705 I AT	1447	15839 AT	1500	17019 S AT
1391	14733_S_AT	1448	15842 AT	1501	17029 S AT
1392	14735 S AT	1449	15857 S AT	1502	17041 S AT
1393	14779 AT	1450	15859 AT	1503	17047_S_AT
1394	14784 AT	1451	15880 AT	1504	17066 S AT
1395	14923 AT	1452	15886 AT	1505	17085 S AT
1396	14947 AT	1453	15906 S AT	1506	17089 S AT
1397	14950 AT	1454	15910 AT -	1507	17179 AT
1398	14990 AT	1455	15937 AT	1508	17180 AT
1399	14998 AT	1456	15957 AT	1509	17228 AT
1400	15005 S AT	1457	15970 S AT	1510	17252 AT
1401	15018 AT	1458	15985 AT	1511	17317 AT
1402	15045 AT	1459	16010 S AT	1512	17338 AT
1403	15046 S AT		16011 S AT	1513	17384 AT
1404	15052 AT		17078 S AT	1514	17387 S AT
1405	15058 S AT	1460	16021 S AT	1515	17400 S AT
1406	15064 AT	1461	16031 AT	1516	17407 S AT
1407	15088 S AT	1462	16038 S AT	1517	17408 AT
1408	15098 S AT	1463	16045 S AT	1518	17413 S AT
1409	15103 S AT	1464	16046 S AT	1519	17416 AT
1410	15109 S AT	1465	16048 AT	1520	17425 S AT
1411	15124_S_AT	1466	16061 S AT	1521	17440 I AT
1412	15127 S AT	1467	16082 S AT	1522	17442 I AT
1413	15145 S AT	1468	16111 F AT	1523	17473_AT
1414	15154_S_AT	1469	16115_S_AT	1524	17484_AT
1415	15161_S_AT	1470	16141_S_AT	1525	17514 S_AT
1416	15189_S_AT	1471	16144_S_AT	1526	17520_S_AT
1417	15214_S_AT	1472	16163_S_AT	1527	17533_S_AT
1418	15255_AT	1473	16173_S_AT	1528	17548_S_AT
1419	15356_AT	1474	16229_AT		19614_AT
1420	15357_AT	1475	16298_AT	1529	17549_S_AT
1421	15364_AT	1476	16301_S_AT	1530	17555_S_AT
1422	15392_AT	1477	16322_AT	1531	17567_AT
1423	15403_S_AT	1478	16342_AT	1532	17654_AT
1424	15437_AT	1479	16351_AT	1533	1 7693_A T
1425	15451_AT	1480	16412_S_AT	1534	17697_AT
1426	15476_AT	1481	16422_AT	1535	17722_AT
1427	15482_AT	1482	16427_AT	1536	17752_AT
1428	15483_S_AT	1483	16438_AT	1537	17755_AT
1429	15521_S_AT	1484	16474_S_AT	1538	17775_AT
1430	15522_I_AT	1485	16482_S_AT	1539	17832_S_AT
1431	15531_I_AT	1486	16485_S_AT	1540	17840_S_AT
1432	15573_AT	1.40=	18052_S_AT	1541	17843_S_AT
1433	15581_S_AT	1487	16493_AT	1542	17855_AT
1434 1435	15586_S_AT 15594_S_AT	1488 1489	16534_S_AT 16555_S_AT	1543 1544	17860_AT 17869_AT
		1489		1544	17888 AT
1436 1437	15609_S_AT 15611_S_AT	1490	16561_S_AT	1545	17888_A1 17899 AT
1437	15611_S_A1 15621_F_AT	1491	17572_S_AT 16592_S_AT	1547	17929_A1 17929_S_AT
1438	15623_F_AT	1491	16615 S AT	1548	17930 S AT
1440	15669 S AT	1492	16637 S AT	1549	17930_3_AT 17932_S_AT
1441	15695 S AT	1494	16692 AT	1550	17936 S AT
1442	15702 S AT	1495	16712 AT	1550	18670 G AT
1772	10.02_5_111	1.173			

TABLE 24 (cont)

			` '		
1551	17957 AT	1606	19152 AT	1663	20040 AT
1552	17961 AT	1607	19156 S AT	1664	20042 S AT
1553	17962 AT	1608	19182 AT	1665	20060 AT
1554	17963 AT	1609	19186 S AT	1005	20438 AT
1555	17971 S AT	1610	19214 AT	1666	20089 AT
1556	17975 AT	1611	19216 AT	1667	20118 AT
1000	18742 F AT	1612	19227 AT	1668	20144 AT
1557	18016 R AT	1613	19243 AT	1669	20149 AT
1558	18069 AT	1614	19288 AT	1670	20179 AT
1559	18122 AT	1615	19359 S AT	1671	20190 AT
1560	18140 AT	1616	19368 AT	1672	20194 AT
1561	18199 AT	1617	19379 AT	1673	20219 AT
1562	18224 S AT	1618	19380_S_AT	1674	20245 S AT
1563	18225 AT	1619	19398 AT	1675	20263 AT
1564	18235 AT	1620	19421 AT	1676	20308 S AT
1565	18259 S AT	1621	19424 AT	1677	20335 S AT
1566	18265 AT	1622	19429 AT	1678	20338 AT
1567	18270 AT1568	1623	19430 AT	1679	20345 AT
	18280 AT	1624	19450 AT	1680	20365 S AT
1569	18289 AT	1625	19457 AT	1681	20382 S AT
1570	18296 AT	1626	19467 AT	1682	20390 S AT
1571	18298 AT	1627	19516 AT	1683	20395 AT
1572	18314 I AT	1628	19545 AT	1684	20420 AT
1573	18318 AT	1629	19564 AT	1685	20421 AT
1574	18325 AT	1630	19577 AT	1686	20432 AT
1575	18351_S_AT	1631	19593 AT	1687	20437 AT
1576	18471 AT	1632	19602 AT	1688	20442 I AT
1577	18482_S_AT	1633	19618 AT	1689	20463_S_AT
1578	18484 AT	1634	19638 AT	1690	20491 AT
1579	18560 AT	1635	19640 AT	1691	20537 AT
1580	18564 AT	1636	19646_S_AT	1692	20573_AT
1581	18590 AT	1637	19656_S_AT	1693	20636_AT
1582	18594_AT	1638	19670_AT	1694	20638_AT
1583	18595_AT	1639	19696_AT	1695	20641_AT
1584	18596_AT	1640	19713_AT	1696	20658_S_AT
1585	18629_S_AT	1641	19 718_A T	1697	20689_S_AT
1586	18637_AT	1642	19722_S_AT	1698	20698_S_AT
1587	18661_AT	1643	19749_AT		
1588	18668_AT	1644	19755_AT		
1589	18699_I_AT	1645	19762_AT		
1590	18747_F_AT	1646	19789_S_AT		
	18789_AT	1647	19815_AT		
1591	18761_AT	1648	19843_AT		
1592	18833_AT	1649	19869_AT		
1593	18875_S_AT	1650	19878_AT		
1594	18894_AT	1651	19883_AT		
1595	18936_AT	1652	19894_AT		
1596	18946_AT	1653	19926_AT		
1597	18953_AT	1654	19944_AT		
1598	18955_AT	1655	19968_AT		
1599	18972_AT	1656	19977_AT		
1600 1601	19008_S_AT	1657 1658	19982_AT 19987_AT		
1601	19108_AT 19123_AT	1659	19987_A1 19991_AT		
1602		1660	20015 AT		
1603	19135_AT 19137_AT	1661	20015_AT 20017_AT		
1604	1913/_AT 19141_AT	1662	20017_A1 20031_AT		
1003	17171_A1	1002	20031_A1		

199 TABLE 25: 2X UP IN COLD, SALT & MANNITOL

19640_at 19646_s_at 19656_s_at 19701_s_at 19843_at 19944_at 19982_at 19997_at 20042_s_at 20042_s_at 20118_at 20144_at 20149_at 20179_at

20194_at

20245_s_at

20390_s_at 20437_at

20463_s_at 20491_at

20658_s_at

20641 at

	TABLE 25:	2X UP IN C
12023_s_at	14733 s_at	17047 s at
12332 s at	14923 at	17179 at
12530 at	14990_at	17180_at
12536_s_at	15005 s at	17252_at
12574 at	15018 at	17384 at
12595 at	15052 at	17407 s at
12698_at	15088_s_at	17484_at
12749 at	15098 s at	17520 s at
12765 at	15103 s at	17555 s at
12769 at	15145 s at	17572 s at
12785_at	15154_s_at	17722_at
12857_at	15161_s_at	17752_at
12964 at	15214 s at	17840_s_at
12972 at	15356 at	17843 s at
12972_at	15521 s at	17860 at
13004 at	15573_at	17000_at
		17929_s_at
13025_at	15586_s_at	
13036_at	15609_s_at	17962_at
13099_s_at	15611_s_at	18052_s_at
13136_at	15621_f_at	18069_at
13176_at	15669_s_at	18122_at
13220_s_at	15695_s_at	18199_at
13225_s_at	15753_at	18259_s_at
13230_s_at	15761_at	18280_at
13239_s_at	15857_s_at	18289_at
13426_at	15871_s_at	18314_i_at
13474_at	15964_s_at	18318_at
13548_at	15970_s_at	18325_at
13555_at	15974_g_at	18482_s_at
13595_at	15997_s_at	18590_at
13627_at	16011_s_at	18594_at
13645_at	16021_s_at	18595_at
13647_at	16038_s_at	18596_at
13706_s_at	16046_s_at	18629_s_at
13965_s_at	16082_s_at	18661_at
13967_at	16111_f_at	18668_at
14080_at	16115_s_at	18699_i_at
14090_i_at	16127_s_at	18722_s_at
14097_at	16141_s_at	18936_at
14116_at	16144_s_at	18953_at
14151_at	16163_s_at	18955_at
14172_at	16236_g_at	18972 at
14192 at	16301 s at	19008_s_at
14244_s_at	16322 at	19152 at
14245_at	16422_at	19186_s_at
14367 at	16474 s at	19214 at
14398 s at	16482 s at	19368_at
14582 at	16485 s at	19379 at
14614_at	16555 s at	19380 s at
14644 s at	16561 s at	19421 at
14645 s at	16592 s at	19545 at
14658 s at	16637_s_at	19614 at
14659 s_at	17041 s at	19638 at
5_6		.000_at

200 TABLE 26: 2X DOWN IN COLD, MANNITOL & SALT, ONLY

	TABLE	26: 2X DOWN IN 6	COLD, MANNITO
12078_at	15189_s_at	17869_at	20015 at
12115_at	15357 at	17888 at	20040_at
12118 at	15364 at	17930 s at	20089 at
12150 at	15403 s at	17932 s. at	20190 at
12271 s at	15476 at	17957 at	20219 at
12276 at	15483 s at	17963 at	20263 at
12338 at	15522 i at	17971 s at	20301 s at
12400 at	15531 i at	17975 at	20308 s at
12430 at	15594 s at	17888_at 17930_s_at 17932_s_at 17957_at 17963_at 17975_at 18016_r_at 18140_at 18224_s_at 18225_at 18225_at 18225_at	20338 at
12538 at	15702 c at	18140 at	20335_at
12622 at	15778 at	18224 e at	20345_at
12630 at	15839 at	18225 at	20442_i_at
12702 c at	15842 at	18229 at	2044Z_1_at
12805 e at	15850 at	18235 at	20537_at
12883 s at	15872 at	18265_at	20636 at
12000_5_at	15072_at	10203_at	20030_at
12932_s_at	15872_at 15880_at 15886_at	18265_at 18270_at 18296_at 18298_at 18471_at	20030_at
		10290_at	20698_s_at
13159_at	15906_S_at	10290_at	
13135_at	15957_at	1047 I_at	
13217_5_dt	10905_at	10004_at	
13279_5_81	10045_s_at	18637_at	
13202_S_at	16061_S_at	18742_T_at	
13432_at	101/3_s_at	18761_at	
13511_at	16298_at	18833_at	
13546_at	15957_at 15985_at 16045_s_at 16045_s_at 16073_s_at 16298_at 16351_at 16412_s_at 16438_at 16493_at 16534_s_at	18295_at 18471_at 18564_at 18637_at 18637_at 18742_fat 18742_fat 18785_s_at 18894_at 18894_at 18926_at 19243_at 19243_at 19243_at 19243_at 19248_at 19388_at 19487_at 19457_at 19467_at 19564_at 19577_at 19564_at 19560_at 19564_at 19560_at 19560_at 19560_at 19600_at 19722_s_at 19722_s_at 18603_at 19600_at 19722_s_at 19722_s_at 18603_at 19722_s_at 19722_s_at 18603_at 19722_s_at 18603_at 19722_s_at 19722_s_at 18603_at 19722_s_at 18603_at 19722_s_at 18603_at 19722_s_at 18603_at 18603_at 19603_at 19722_s_at 18603_at 18603_at 19603_at 19603_at 19722_s_at 18603_at 18603_a	
13547_S_at	10412_S_at	18894_at	
13610 c ot	16436_at	10946_at	
13640 of	10493_at	19123_at	
13775 of	10534_S_at	19216_at	
13771 at	10009_S_at	19243_at	
13016 ot	10013_s_at	19207_S_at	
14028 at	16790 of	19200_at	
14020_at	16709_at	19390_at	
14005_at	16071 c of	19424_at	
14040_at	17019 c at	19430_at	
14077_at	17010_s_at	19450_at	
14170 at	17029_s_at	19457_at	
1/170_at	17009_3_at	19407_at	
14248 at	17220_at	19510_at	
14381 at	17387 s at	19504_at	
14301_at	17307_S_dt	19577_at	
14304_at	17415_5_dt	19595_at	
14507_at	17410_at	19002_at	
14705 i at	17425_s_at 17440_i_at 17473_at	19010_at	
14709_r_at	17440_1_at	19070_at	
14779 at	17522 c of	19090_at	
14947_at	17533_s_at 17549_s_at 17654_at	19722_s_at	
14950 at	17654 at	10755 at	
14998 at	17693 at	19755_at	
15045 at	17693_at 17697_at	19926_at	
15109_s_at	17755_at	19968_at	
15109_s_at	17733_at 17832_s_at		
10124_5_all	1/032_S_8[19977_at	

201 TABLE 27: 2X ROOT SPECIFIC (COLD, SALINE & OSMOTIC STRESSES)

		OOI SI ECIFIC (CO)	D, SALINE & OSMOTIC S
11997_at	14069_at	16052_at	18327_s_at
12004_at	14072_at	16053_i_at	18597_at
12051_at	14073_at	16105_s_at	18607_s_at
12072_at	14097_at	16161 s at	18636 at
12150_at	14139_at	16165 s at	18663 s at
12151_at	14235_at	16298_at	18782_at
12166_i_at	14250_r_at	16334_s_at	18885_at
12219_at	14578_s_at	16422 at	18888_at
12315_at	14582_at	16427 at	18942 at
12332 s at	14640 s at	16440_s_at	18955 at
12374 i at	14643 s at	16442 s at	19060_at
12482_s at	14644 s at	16468 at	19108 at
12515 at	14658 s at	16488_at	19135 at
12522 at	14659 s at	16511 at	19137 at
12538 at	14711 s at	16529 at	19195 at
12571_s at	14900 at	16553 f at	19263 at
12574 at	14924 at	16568_s_at	19376_at
12609 at	14990 at	16914 s at	19406 at
12678 i at	15018_at	16965 s at	19432 s at
12698_at	15022 at	16981_s_at	19835 at
12749 at	15107 s at	16989 at	19836_at
12760 g at	15116_f_at	17033 s at	19840 s at
12765 at	15120 s at	17066_s_at	19841 at
12768_at	15124 s at	17085_s_at	19843_at
12769 at	15131 s at	17252 at	19926 at
12772_at	15132_s_at	17376 at	19972_at
12777_i_at	15137_s_at	17378 at	19977_at
12958 at	15184 s at	17388_at	19991_at
12989 s at	15188 s at	17415 at	20034 i at
13015_s_at	15208_s_at	17429 s at	20042 s at
13134 s at	15252_g_at	17463 at	20189 at
13146_s_at	15343_at	17485_s_at	20194 at
13172_s_at	15389 at	17490 s at	20200 at
13178_at	15392 at	17567_at	20214 i at
13179_at	15448 at	17585_s_at	20239_g_at
13187_i_at	15503 at	17595 s at	20262 at
13211_s_at	15531_i_at	17840_s_at	20269 at
13239_s_at	15594 s at	17860 at	20294 at
13273_s_at	15609 s at	17880 s at	20312 s at
13297_s_at	15623_f_at	17894_at	20382_s_at
13549_at	15639_s_at	17896 at	20396 at
13604_at	15670_s_at	17899_at	20432 at
13629_s_at	15680_s_at	17911_at	20444_at
13706_s_at	15859_at	17935 at	20446 s at
13714_at	15900_at	17961_at	20480 s at
13751_at	15923_at	18024 s at	20586_i_at
13895_at	15962_s_at	18122_at	20612_s_at
13933_at	15964_s_at	18222_at	20672_at
13967_at	15965_at	18224_s_at	20686_at
13985_s_at	15975_s_at	18252_at	20689_s_at
14028_at	15985_at	18255_at	
14030_at	16001_at	18269_s_at	
14058_at	16048_at	18270_at	

202
TABLE 28: 2X LEAF SPECIFIC (COLD, SALINE & OSMOTIC STRESSES)

IABLE
16136_s_a
16172_s_a
16316_at
16385_s_a
16455_at
16485_s_a
16512_s_a
16547_s_a
16548_s_a
16629_s_a
16673_at
16899_at
17010_s_a
17018_s_a
17054_s_a
17095_s_at
17097_s_a
17273_at
17394_s_at
17420_at
17449_s_at
17600_s_at
17843_s_at
17913_s_at
17966_at
18003_at
18081_at
18560_at
18588_at
18626_at 18644_at
18666_s_at 18742 f at
18742_f_at
18994_at
19227_at
19373 at
19834 at
19867 at
19998 at
20062 at
20199 at
20155_at
20284 at
20437 at
20442 i at
20450 at
20468_at
20547 at
20635 s at

15638_s_at

19755 at

CRIPTION (COLD, SALINE & OSMOTIC STRESSES)

	TABI E 20. 2V T	203 RANSCRIPTION (Co
12000 -1		
12068_at	15665_s_at 15679_s_at	19836_at
12 100_1_at	15679_s_at	19860_at
12374_I_8t	15720_at	19866_at
12166 i_at 12374_i_at 12392_at 12431_at	15871_s_at 16072_s_at	19898_at
12431_at	160/2_s_at	20262_at
12450_s_at 12503_at	160/3_f_at	20335_s_at
12503_at 12536_s_at 12540_s_at 12541_at	16073_f_at 16105_s_at 16111_f_at 16127_s_at 16534_s_at	20362_at
12536_s_at	16111_1_at	20424_at
12540_s_at	1612/_s_at	20437_at
12541_at	16534_s_at	20456_at
12541_at 12587_at 12594_at 12595_at	16582_s_at 16589_s_at	20515_s_at
12594_at	16589_s_at	20635_s_at
12595_at	16747_at	
12704_f_at 12705_f_at	17019_s_at	
12705_f_at	17129_s_at	
12709_f_at 12712_f_at	17160_at	
12/12_f_at	17520_s_at	
12719 f at 12724 f at 12725 r at	17538_s_at	
12/24_f_at	17555_s_at	
12725_F_at	17609_at	
12/26_f_at	17896_at	
12734_I_at	17971_s_at	
12/30_1_at	17975_at	
12726_f_at 12726_f_at 12734_f_at 12736_f_at 12737_f_at 12812_at 12949_at 12951_at	17978_s_at	
12012_at	18121_s_at	
12949_at	18167_s_at	
12951_at 12966_s_at	18197_at	
13023_at 13034_s_at 13087_at 13270_at	18576 c ot	
13034_S_at	10070_S_at	
13007_at	19729_S_at	
13273 c at	18742 f at	
13432 at	18744 f at	
13555 at	18745 f at	
13688 s at	18747 f at	
13270_at 13273_s_at 13432_at 13555_at 13688_s_at 13714_at	18750 f at	
13965 s at	18751 f at	
13965_s_at 13987_s_at 14003_at 14144_at	18750_f_at 18751_f_at 18789_at	
14003 at	18834_at	
14144 at	18942 at	
14178 at	19083_at	
14178_at 14223_at 14235_at	19202_at	
14235 at	19209_s_at	
14303_s_at	19232_s_at	
14393_at	19315_at	
14553 at	19489_s_at	
14781_at	19611 s at	
15046 s at	19646_s_at	
15053 s at		
15053_s_at 15214_s_at 15510 r_at	19707_s_at 19722_s_at 19744_at	
15510 rat	19744_at	
15620	40755 - I	

13192 s at 13193 s at 13587_at 13995 at 14335_at 15073 at 15171_s_at 15240 at 15586_s_at 15641_s_at 15651 f at 15990 at 16232 s at 16576 f at 16753_at 17423_s_at 17525 s at 17537 s at 17929_s_at 17954_s_at 18012_s_at 18308_i_at 18616 at 18847 at 18936 at 18980 at 19243_at 19263 at 19638 at 19883 at 19932 at 20333 at 20393 at

20570 at

12470_at 12556_at 13128_at 13135_s_at 13180_s_at

TABLE 31: 2X KINASES (COLD, SALINE & OSMOTIC STRESSES)

20144_at 20219_at 20223_at 20232_s_at 20235_i_at 20282_s_at 20298_at 20396_at 20439_at 20462_at

	TABLE 3
12253_g_at	16059_s_at
12270_at	16087_s_at
12271 s at	16088_f_at
12276_at	16125 s at
12278 at	16137_s_at
12284_at	16140 s at
12300 at	16143 s at
12307 at	16144 s at
12353 at	16160 f at
12357 s at	16171 s at
12390 at	16357 at
12394 at	16412 s at
12395 s at	16568 s at
12408_at	16570 s at
12452 at	16571 s at
12477_at	16584 s at
12490 at	16651 s at
12497 at	16652 s at
12532 at	16672 at
12697_at	16818 s at
12901 s at	16840 at
12902 at	17068_s_at
12958 at	17122_s_at
12959 at	17252 at
13068 at	17323 at
13246 at	17475 at
13324 at	17752 at
13332_at	17921_s_at
13362 s at	17933_at
13370_at	17935 at
13550_at	18013_r_at
14030_at	18046 s at
14048_at	18122 at
14194_at	18176 at
14196_at	18316_at
14217_at	18455_at
14459_at	18459_at
14603_at	18482_s_at
14637_s_at	18543_at
14686_s_at	18706_s_at
15005_s_at	18782_at
15175_s_at	18924_at
15270_at	19117 s at
15475_s_at	19437_s_at
15497_s_at	19442_at
15577_s_at	19458_at
15616_s_at	19464_at
15633_s_at	19469_at
15634_s_at	19562_at
15668_s_at	19655_at
15680_s_at	19749_at
15798_at	19854_at
16034_at	19904_at